PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:

C12Q 1/70

A2

(11) International Publication Number: WO 99/67428

(43) International Publication Date: 29 December 1999 (29.12.99)

EP

(21) International Application Number: PCT/EP99/04317

(22) International Filing Date: 22 June 1999 (22.06.99)

(30) Priority Data: 98870143.9 24 June 1998 (24.06.98)

(71) Applicant (for all designated States except US): INNOGENET-ICS N.V. [BE/BE]; Industriepark Zwijnaarde 7, P.O. Box 4, B-9052 Ghent (BE).

(72) Inventor; and
(75) Inventor/Applicant (for US only): STUYVER, Lieven

[BE/BE]; Holestraat 8, B-9552 Herzele (BE).

(74) Common Representative: INNOGENETICS N.V.; Industriepark Zwijnaarde 7, P.O. Box 4, B-9052 Ghent (BE).

(81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published

Without international search report and to be republished upon receipt of that report.

(54) Title: METHOD FOR DETECTION OF DRUG-SELECTED MUTATIONS IN THE HIV PROTEASE GENE

(57) Abstract

The present invention relates to a method for the rapid and reliable detection of drug-selected mutations in the HIV protease gene allowing the simultaneous characterization of a range of codons involved in drug resistance using specific sets of probes optimized to function together in a reverse-hybridization assay. More particularly, the present invention relates to a method for determining the susceptibility to antiviral drugs of HIV viruses in a biological sample, with said method comprising: a) if need be, releasing, isolating or concentrating the polynucleic acids present in the sample; b) if need be amplifying the relevant part of the protease gene of HIV with at least one suitable primer pair; c) hybrydizing the polynucleic acids of step a) or b) with at least one of the following probes: probes specifically hybridizing to a target sequence comprising codon 30; probes specifically hybridizing to a target sequence comprising codon 46 and/or 48; probes specifically hybridizing to a target sequence comprising codon 50; probes specifically hybridizing to a target sequence comprising codon 51; probes specifically hybridizing to a target sequence comprising codon 90; or the complement of said probes; further characterized in that said probes specifically hybridize to any of the target sequences presented in figure (1), or the complement of said target sequences; d) inferring from the result of step c) whether or not a mutation giving rise to drug resistance is present in any of said target sequences.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	l.S	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria .	FR	France	LU	Luxembourg	SN	Senegal
ΑU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
ΑZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	· MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Treland	MN	Mongolia	UA.	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	ıc	Saint Lucia	RU	Russian Federation		
DE	Germany	L	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

WO 99/67428 PCT/EP99/04317

METHOD FOR DETECTION OF DRUG-SELECTED MUTATIONS IN THE HIV PROTEASE GENE

5

10

15

20

25

30

35

1. FIELD OF THE INVENTION

The present invention relates to the field of HIV diagnosis. More particularly, the present invention relates to the field of diagnosing the susceptibility of an HIV sample to antiviral drugs used to treat HIV infection.

The present invention relates to a method for the rapid and reliable detection of drug-selected mutations in the HIV protease gene allowing the simultaneous characterization of a range of codons involved in drug resistance using specific sets of probes optimized to function together in a reverse-hybridization assay.

2. BACKGROUND OF THE INVENTION

The human immunodeficiency virus (HIV) is the ethiological agent for the acquired immunodeficiency syndrome (AIDS). HIV, like other retroviruses, encodes an aspartic protease that mediates the maturation of the newly produced viral particle by cleaving viral polypeptides into their functional forms (Hunter et al). The HIV protease is a dimeric molecule consisting of two identical subunits each contributing a catalytic aspartic residue (Navia et al, Whodawer et al, Meek et al). Inhibition of this enzyme gives rise to noninfectious viral particles that cannot establish new cycles of viral replication (Kohl et al, Peng et al).

Attempts to develop inhibitors of HIV-1 protease were initially based on designing peptide compounds that mimicked the natural substrate. The availability of the 3-dimensional structure of the enzyme have more recently allowed the rational design of protease inhibitors (PI) using computer modeling (Huff et al, Whodawer et al). A number of second generation PI that are partially peptidic or entirely nonpeptidic have proven to exhibit particularly potent antiviral effects in cell culture. Combinations of various protease inhibitors with nucleoside and non-nucleoside RT inhibitors have also been studied extensively in vitro. In every instance, the combinations have been at least additive and usually synergistic.

In spite of the antiviral potency of many recently developed HIV-1 PI, the emergence of virus variants with decreased sensitivity to these compounds has been described both in cell culture and in treated patients thereby escaping the inhibitory effect of the antiviral (Condra et al.). Emergence of

10

15

20

25

30

35

resistant variants depends on the selective pressure applied to the viral population. In the case of a relatively ineffective drug, selective pressure is low because replication of both wild-type virus and any variants can continue. If a more effective drug suppresses replication of virus except for a resistant variant, then that variant will be selected. Virus variants that arise from selection by PI carry several distinct mutations in the protease coding sequence that appear to emerge sequentially. A number of these cluster near the active site of the enzyme while others are found at distant sites. This suggests conformational adaptation to primary changes in the active site and in this respect certain mutations that increase resistance to PI also decrease protease activity and virus replication.

Amongst the PI, the antiviral activity of the PI ritonavir (A-75925; ABT-538), nelfinavir (AG-1343), indinavir (MK-639; L735; L524) and saquinavir (Ro 31-8959) have been approved by the Food and Drug Administration and are currently under evaluation in clinical trials involving HIV-infected patients. The VX-487 (141W94) antiviral compound is not yet approved. The most important mutations selected for the above compounds and leading to gradually increasing resistance are found at amino acid (aa) positions 30 (D to N), 46 (M to I), 48 (G to V), 50 (I to V), 54 (I to A, I to V), 82 (V to A, or F, or I, or T), 84 (I to V) and 90 (L to M). Other mutations associated with drug resistance to the mentioned compounds have been described (Schinazi et al). Saquinavir-resistant variants, which usually carry mutations at amino acid positions 90 and/or 48, emerge in approximately 45% of patients after 1 year of monotherapy. Resistance appears to develop less frequently with higher doses of saquinavir. Resistance to indinavir and ritonavir requires multiple mutations; usually at greater than 3 and up to 11 sites, with more amino acid substitutions conferring higher levels of resistance. Resistant isolates usually carry mutations at codons 82, 84, or 90. In the case of ritonavir, the mutation at codon 82 appears first in most patients. Although mutant virions resistant to saquinavir are not cross-resistant to indinavir or ritonavir, isolates resistant to indinavir are generally ritonavir resistant and visa versa. Resistance to either indinavir or ritonavir usually results in cross-resistance to saquinavir. Approximately one third of indinavir resistant isolates are cross-resistant to nelfinavir as well.

The regime for an efficient antiviral treatment is currently not clear at all. Patterns of reduced susceptibility to HIV protease inhibitors have been investigated *in vitro* by cultivating virus in the presence of PI. These data, however, do not completely predict the pattern of amino-acid changes actually seen in patients receiving PI. Knowledge of the resistance and cross-resistance patterns should facilitate selection of optimal drug combinations and selection of sequences with non-overlapping resistance patterns. This would delay the emergence of cross-resistant viral strains and prolong the duration of effective antiretroviral activity in patients. Therefore, there is need for methods and systems that detect these mutational events in order to give a better insight into the mechanisms of HIV resistance. Further, there is need for methods and systems which can provide data important for the antiviral therapy to follow in a more time-efficient and economical manner compared to the conventional cell-culture selection techniques.

10

15

20

25

30

35

3. AIMS OF THE INVENTION

It is an aim of the present invention to develop a rapid and reliable detection method for determination of the antiviral drug resistance of viruses, which contain protease genes such as HIV retroviruses present in a biological sample.

More particularly it is an aim of the present invention to provide a genotyping assay allowing the detection of the different HIV protease gene wild type and mutation codons involved in the antiviral resistance in one single experiment.

It is also an aim of the present invention to provide an HIV protease genotyping assay or method which allows to infer the nucleotide sequence at codons of interest and/or the amino acids at the codons of interest and/or the antiviral drug selected spectrum, and possibly also infer the HIV type or subtype isolate involved.

Even more particularly it is an aim of the present invention to provide a genotyping assay allowing the detection of the different HIV protease gene polymorphisms representing wild-type and mutation codons in one single experimental setup.

It is another aim of the present invention to select particular probes able to discriminate wild-type HIV protease sequences from mutated or polymorphic HIV protease sequences conferring resistance to one or more antiviral drugs, such as ritonavir (A-75925; ABT-538), nelfinavir (AG-1343), indinavir (MK-639; L735; L524), saquinavir (Ro 31-8959) and VX-478 (141W94) or others (Shinazi *et al*).

It is more particularly an aim of the present invention to select particular probes able to discriminate wild-type HIV protease sequences from mutated or polymorphic HIV protease sequences conferring resistance to ritonavir (A-75925; ABT-538).

It is more particularly an aim of the present invention to select particular probes able to discriminate wild-type HIV protease sequences from mutated HIV protease sequences conferring resistance to nelfinavir (AG-1343).

It is more particularly an aim of the present invention to select particular probes able to discriminate wild-type HIV protease sequences from mutated HIV protease sequences conferring resistance to indinavir (MK-639; L735; L524).

It is more particularly an aim of the present invention to select particular probes able to discriminate wild-type HIV protease sequences from mutated HIV protease sequences conferring resistance to saquinavir (Ro 31-8959).

It is more particularly an aim of the present invention to select particular probes able to discriminate wild-type HIV protease sequences from mutated HIV protease sequences conferring resistance to VX-478 (141W94).

It is also an aim of the present invention to select particular probes able to determine and/or infer

10

15

20

25

30

cross-resistance to HIV protease inhibitors.

It is more particularly an aim of the present invention to select particular probes able to discriminate wild-type HIV protease from mutated HIV protease sequences involving at least one of amino acid positions 30 (D to N), 46 (M to I), 48 (G to V), 50 (I to V), 54 (I to A or V), 82 (V to A or F or I or T), 84(I to V) and 90 (L to M) of the viral protease gene.

It is particularly an aim of the present invention to select a particular set of probes, able to discriminate wild-type HIV protease sequences from mutated HIV protease sequences conferring resistance to any of the antiviral drugs defined above with this particular set of probes being used in a reverse hybridization assay.

It is moreover an aim of the present invention to combine a set of selected probes able to discriminate wild-type HIV protease sequences from mutated HIV protease sequences conferring resistance to antiviral drugs with another set of selected probes able to identify the HIV isolate, type or subtype present in the biological sample, whereby all probes can be used under the same hybridization and wash-conditions.

It is also an aim of the present invention to select primers enabling the amplification of the gene fragment(s) determining the antiviral drug resistance trait of interest.

It is also an aim of the present invention to select particular probes able to identify mutated HIV protease sequences resulting in cross-resistance to antiviral drugs.

The present invention also aims at diagnostic kits comprising said probes useful for developing such a genotyping assay.

The present invention also aims at diagnostic kits comprising said primers useful for developing such a genotyping assay.

4. DETAILED DESCRIPTION OF THE INVENTION.

All the aims of the present invention have been met by the following specific embodiments.

According to one embodiment, the present invention relates to a method for determining the susceptibility to antiviral drugs of HIV viruses in a biological sample, with said method comprising:

- a) if need be, releasing, isolating or concentrating the polynucleic acids present in the sample;
- b) if need be amplifying the relevant part of the protease gene of HIV with at least one suitable primer pair;
- c) hybridizing the polynucleic acids of step a) or b) with at least one of the following probes:

 probes specifically hybridizing to a target sequence comprising codon 30;
- probes specifically hybridizing to a target sequence comprising codon 46 and/or 48; probes specifically hybridizing to a target sequence comprising codon 50;

10

15

20

25

probes specifically hybridizing to a target sequence comprising codon 54; probes specifically hybridizing to a target sequence comprising codon 82 and/or 84; probes specifically hybridizing to a target sequence comprising codon 90; or the complement of said probes,

further characterized in that said probes specifically hybridize to any of the target sequences presented in figure 1, or to the complement of said target sequences;

d) inferring from the result of step c) whether or not a mutation giving rise to drug resistance is present in any of said target sequences.

The numbering of HIV-1 protease gene encoded amino acids is as generally accepted in literature.

Mutations that give rise to an amino acid change at position 48 or 90 are known to confer resistance to saquinavir (Erlebe et al; Tisdale et al). An amino acid change at codon 46 or 54 or 82 or 84 results in ritonavir and indinavir resistance (Kempf et al; Emini et al; Condra et al). Amino acid changes at positions 30 and 46 confer resistance to nelfinavir (Patick et al) and amino acid changes at position 50 confers resistance to VX-487 (Rao et al). Therefore, the method described above allows to determine whether a HIV strain is susceptible or resistant to any of the drugs mentioned above. This method can be used, for instance, to screen for mutations conferring resistance to any of the mentioned drugs before initiating therapy. This method may also be used to screen for mutations that may arise during the course of therapy (i.e. monitoring of drug therapy). It is obvious that this method may also be used to determine resistance to drugs other than the above-mentioned drugs, provided that resistance to these other drugs is linked to mutations that can be detected by use of this method. This method may also be used for the specific detection of polymorphic nucleotides. It is to be understood that the said probes may only partly overlap with the targets sequences of figure 1, table 2 and table 3, as long as they allow for specific detection of the relevant polymorphic nucleotides as indicated above. The sequences of figure 1, table 2 and table 3 were derived from polynucleic acid fragments comprising the protease gene. These fragments were obtained by PCR amplification and were inserted into a cloning vector and sequence analyzed as described in example 1. It is to be noted that some polynucleic acid fragments comprised polymorphic nucleotides in their sequences, which have not been previously disclosed. These novel polymorphic nucleotide sequences are represented in table 4 below.

30 TABLE 4: Polymorphic nucleotide sequences.

	51	5 2	53	54	55	56	57	58	codon position		
	gga	ggt	ttt	atc	aaa	gta	aga	cag	consensus	sequence	
	GGA	GGT	TTT	ATC	AAA	GTC	AGA	CAA	SEQ ID NO	478	
35	GGA	GGT	TTC	ATT	AAG	GTA	AAA	CAG	SEQ ID NO	479	
	GGA	GGT	TTT	ATT	AAG	GTA	AGA	CAG	SEQ ID NO	480	

GGA GGT TTT ATT AAA GTA AGA CAA

6

SEQ ID NO 481

	GGA	GGC	TTT	ATC	AAA	GTA	AGA	CAA		SEQ ID NO	482
	GGA	ggţ	TTT	ATC	AAA	GTC	AGA	CAA		SEQ ID NO	483
5	78	79	80	81	82	83	84	85		codon pos	ition
	gga	cct	aca	cct	gtc	aac	ata	att	gg	consensus	sequence
	GGA	CCT	ACA	CCG	GTC	AAC	ATA	ATT	GG	SEQ ID NO	484
	GGA	CCT	ACA	CCT	GCC	AAT	ATA	ATT	GG	SEQ ID NO	485
	GGA	CCT	ACG	CCC	TTC	AAC	ATA	ATT	GG	SEQ ID NO	486
10	GGA	CCG	ACA	CCT	GTC	ACC	ATA	ATT	GG	SEQ ID NO	487
	GGA	CCT	ATA	CCT	GTC	AAC	ATA	ATT	GG	SEQ ID NO	488

	6/	88	89	90	ЭT	92	93	94	codo	codon position		
a	aga	aat	ctg	ttg	act	cag	att	ggc	cons	ens	us	sequence
A	AAA	AAT	CTG	ATG	ACT	CAG	ATT	GGC	SEQ	ID	NO	489
A	AGA	ACT	CTG	TTG	ACT	CAG	CTT	GGA	SEQ	ID	NO	490
A	AGA	AAT	ATG	ATG	ACC	CAG	CTT	GGC	SEQ	ID	NO.	491
A	AGA	AAT	ATA	ATG	ACT	CAG	CTT	GGA	SEQ	ID	NO	492
A	AGA	AAT	CTG	CTG	ACT	CAG	TTA	GGG	SEQ	ID	N0	493
A	AGA	TAA	CTG	TTG	ACA	CAG	CTT	GGC	SEQ	ID	N0	494
A	AGA	TAA	ATG	TTG	ACT	CAG	CTT	GGT	SEQ	ID	N0	495
A	AGA	TAA	TTG	TTG	ACT	CAG	ATT	GGG	SEQ	ID	И0	496
A	AGA	AAT	ATG	TTG	ACT	CAG	CTT	GGT	SEQ	ID	NO	497
A	AGA	AAT	ATG	TTG	ACT	CAG	CTT	GGA	SEQ	ID	N0	498
A _.	AGA	AAT	CTG	TTG	ACT	CAG	CTT	GGA	SEQ	ID	NO	499
A	AGA	AAC	CTG	TTG	ACT	CAA	CTT	GGT	SEQ	ID	ио	500

The present invention thus also relates to these novel sequences, or a fragment thereof, wherein said fragment consists of at least 10, preferably 15, even more preferably 20 contiguous nucleotides and contains at least one polymorphic nucleotide. It is furthermore to be understood that these new

10

15

20

25

30

35

polymorphic nucleotides may also be expected to arise in another sequence context than in the mentioned sequences. For instance a G at the third position of codon 55 is shown in SEQ ID N° 478 in combination with a T at the third position of codon 54, but a G at the third position of codon 55 may also be expected to occur in the context of a wild type sequence. It is also to be understood that the above mentioned specifications apply to the complement of the said target sequences as well. This applies also to Figure 1.

According to a preferred embodiment the present invention relates to a method as indicated above, further characterized in that said probes are capable of simultaneously hybridizing to their respective target regions under appropriate hybridization and wash conditions allowing the detection of the hybrids formed.

According to a preferred embodiment, step c is performed using a set of at least 2, preferably at least 3, more preferably at least 4 and most preferably at least 5 probes meticulously designed as such that they show the desired hybridization results. In general this method may be used for any purpose that relies on the presence or absence of mutations that can be detected by this method, e.g. for genotyping. The probes of table 1 have been optimized to give specific hybridization results when used in a LiPA assay (see below), as described in examples 2 and 3. These probes have thus also been optimized to simultaneously hybridize to their respective target regions under the same hybridization and wash conditions allowing the detection of hybrids. The sets of probes for each of the codons 30, 46/48, 50, 54 and 82/84 have been tested experimentally as described in examples 2 and 3. The reactivity of the sets shown in table 1 with 856 serum samples from various geographic origins was evaluated. It was found that the sets of probes for codons 30, 46/48, 50, 54 and 82/84 reacted with 98.9%, 99.6%, 98.5%, 99.2%, 95.4% and 97.2% of the test samples, respectively. The present invention thus also relates to the sets of probes for codons 30, 46/48, 50, 54, 82/84 and 90, shown in table 1 and table 7.

According to another even more preferred embodiment, the present invention relates to a method as defined above, further characterized in that:

step b) comprises amplifying a fragment of the protease gene with at least one 5'-primer specifically hybridizing to a target sequence located between nucleotide position 210 and nucleotide position 260 (codon 87), more preferably between nucleotide position 220 and nucleotide position 260 (codon 87), more preferably between nucleotide position 230 and nucleotide position 260 (codon 87), even more preferably at nucleotide position 241 to nucleotide position 260 (codon 87) in combination with at least one suitable 3'-primer, and

step c) comprises hybridizing the polynucleic acids of step a) or b) with at least one of the probes specifically hybridizing to a target sequence comprising codon 90.

According to another even more preferred embodiment, the present invention relates to a method as defined above, further characterized in that:

10

15

20

25

30

35

step b) comprises amplifying a fragment of the protease gene with at least one 3'-primer specifically hybridizing to a target sequence located between nucleotide position 253 (codon 85) and nucleotide positions 300, more preferably between nucleotide position 253 (codon 85) and nucleotide positions 290, more preferably between nucleotide position 253 (codon 85) and nucleotide positions 280, even more preferably at nucleotide position 253 (codon 85) to nucleotide position 273 (codon 91), in combination with at least one suitable 5'-primer, and step c) comprises hybridizing the polynucleic acids of step a) or b) with at least one of the probes specifically hybridizing to a target sequence comprising any of codons 30, 46, 48, 50, 52, 54, 82 and 84.

It has been found, unexpectedly, that an amplified nucleic acid fragment comprising all of the above-mentioned codons, does not hybridize optimally to probes comprising codon 82, 84 or 90. On the other hand, a shorter fragment, for instance the fragment which is amplified by use of the primers Prot41bio and Prot6bio with respectively seq id no 5 and seq id no 4, hybridizes better to probes comprising codon 90. Better hybridization is also obtained when the fragment is amplified with primer Prot41bio in combination with primers Prot6abio, Prot6bbio, Prot6cbio and Prot6dbio The present invention thus also relates to a method as defined above, further characterized in that the 5'-primer is seq id no 5 and at least one 3' primer is chosen from seq id no 4, seq id no506, seq id no 507, seq id no 508, and seq id no 509. Likewise, another shorter fragment, for instance the fragment which is amplified by use of the primers Prot2bio and Prot31bio with respectively seq id no 3 and seq id no 6, was found to hybridize better to probes comprising codon 82 and/or 84. Hence the present invention also relates to a method as defined above, further characterized in that the 5'-primer is seq id no 5 and at least one 3'-primer is chosen from seq id no 4, seq id no 506, seq id no 507, seq id no 508, and seq id no 509.

New sets of amplification primers as mentioned in example 1 were selected. The present invention thus also relates to primers: prot 16 (SEQ ID NO 501), prot 5 (SEQ ID NO 5), prot2a bio (SEQ ID NO 503), prot2b bio (SEQ ID NO 504), prot31 bio (SEQ ID NO 6), prot41-bio (SEQ ID NO 505), prot6a (SEQ ID NO 506), prot6b (SEQ ID NO 507), prot6c (SEQ ID NO 508) and prot6d (SEQ ID NO 509). A number of these primers are chemically modified (biotinylated), others are not. The present invention relates to any of the primers mentioned, primers containing unmodified nucleotides, or primers containing modified nucleotides.

Different techniques can be applied to perform the sequence-specific hybridization methods of the present invention. These techniques may comprise immobilizing the amplified HIV polynucleic acids on a solid support and performing hybridization with labeled oligonucleotide probes. HIV polynucleic acids may also be immobilized on a solid support without prior amplification and subjected to hybridization. Alternatively, the probes may be immobilized on a solid support and hybridization may be performed with labeled HIV polynucleic acids, preferably after amplification. This technique is called reverse hybridization. A convenient reverse hybridization technique is the line probe assay (LiPA). This

10

15

30

35

assay uses oligonucleotide probes immobilized as parallel lines on a solid support strip (Stuyver et al., 1993). It is to be understood that any other technique based on the above-mentioned methods is also covered by the present invention.

According to another preferred embodiment, the present invention relates to any of the probes mentioned above and/or to any of the primers mentioned above, with said primers and probes being designed for use in a method for determining the susceptibility to antiviral drugs of HIV viruses in a sample. According to an even more preferred embodiment, the present invention relates to the probes with seq id no 7 to seq id no 477 and seq id no510 to seq id no 519, more preferably to the seq id no mentioned in Table 1 and Table 7, and to the primers with seq id no 3, 4, 5 and 6, 501, 502, 503, 504, 505, 506, 507, 508 and 509. The skilled man will recognize that addition or deletion of one or more nucleotides at their extremities may adapt the said probes and primers. Such adaptations may be required if the conditions of amplification or hybridization are changed, or if the amplified material is RNA instead of DNA, as is the case in the NASBA system.

According to another preferred embodiment, the present invention relates to a diagnostic kit enabling a method for determining the susceptibility to antiviral drugs of HIV viruses in a biological sample, with said kit comprising:

- a) when appropriate, a means for releasing, isolating or concentrating the polynucleic acids present in said sample;
- b) when appropriate, at least one of the primers of any of claims 4 to 6;
- 20 c) at least one of the probes of any of claims 1 to 3, possibly fixed to a solid support;
 - d) a hybridization buffer, or components necessary for producing said buffer;
 - e) a wash solution, or components necessary for producing said solution;
 - f) when appropriate, a means for detecting the hybrids resulting from the preceding hybridization;
 - h) when appropriate, a means for attaching said probe to a solid support.

25 DEFINITIONS

The following definitions serve to illustrate the terms and expressions used in the present invention.

The term "antiviral drugs" refers particularly to any antiviral protease inhibitor. Examples of such antiviral drugs and the mutation they may cause in the HIV protease gene are disclosed in Schinazi et al., 1997. The contents of the latter two documents particularly are to be considered as forming part of the present invention. The most important antiviral drugs focussed at in the present invention are disclosed in Tables 1 to 2.

The target material in the samples to be analyzed may either be DNA or RNA, e.g.: genomic DNA, messenger RNA, viral RNA or amplified versions thereof. These molecules are also termed polynucleic acids.

It is possible to use genomic DNA or RNA molecules from HIV samples in the methods according to the present invention.

10

15

20

25

30

35

Well-known extraction and purification procedures are available for the isolation of RNA or DNA from a sample (fi. in Maniatis et al., Molecular Cloning: A Laboratory Manual, 2nd Edition, Cold Spring Harbour Laboratory Press (1989)).

The term "probe" refers to single stranded sequence-specific oligonucleotides, which have a sequence, which is complementary to the target sequence to be detected.

The term "target sequence" as referred to in the present invention describes the wild type nucleotide sequence, or the sequence comprising one or more polymorphic nucleotides of the protease gene to be specifically detected by a probe according to the present invention. This nucleotide sequence may encompass one or several nucleotide changes. Target sequences may refer to single nucleotide positions, codon positions, nucleotides encoding amino acids or to sequences spanning any of the foregoing nucleotide positions. In the present invention said target sequence often includes one or two variable nucleotide positions.

The term "polymorphic nucleotide" indicates a nucleotide in the protease gene of a particular HIV virus that is different from the nucleotide at the corresponding position in at least one other HIV virus. The polymorphic nucleotide may or may not give rise to resistance to an antiviral drug. It is to be understood that the complement of said target sequence is also a suitable target sequence in some cases. The target sequences as defined in the present invention provide sequences which should be complementary to the central part of the probe which is designed to hybridize specifically to said target region.

The term "complementary" as used herein means that the sequence of the single stranded probe is exactly the (inverse) complement of the sequence of the single-stranded target, with the target being defined as the sequence where the mutation to be detected is located.

"Specific hybridization" of a probe to a target sequence of the HIV polynucleic acids means that said probe forms a duplex with part of this region or with the entire region under the experimental conditions used, and that under those conditions said probe does not form a duplex with other regions of the polynucleic acids present in the sample to be analyzed.

Since the current application requires the detection of single basepair mismatches, very stringent conditions for hybridization are required, allowing in principle only hybridization of exactly complementary sequences. However, variations are possible in the length of the probes (see below), and it should be noted that, since the central part of the probe is essential for its hybridization characteristics, possible deviations of the probe sequence versus the target sequence may be allowable towards head and tail of the probe, when longer probe sequences are used. These variations, which may be conceived from the common knowledge in the art, should however always be evaluated experimentally, in order to check if they result in equivalent hybridization characteristics than the exactly complementary probes.

Preferably, the probes of the invention are about 5 to 50 nucleotides long, more preferably from about 10 to 25 nucleotides. Particularly preferred lengths of probes include 10, 11, 12, 13, 14, 15, 16, 17,

10

15

20

25

30

35

18, 19, 20, 21, 22, 23, 24 or 25 nucleotides. The nucleotides as used in the present invention may be ribonucleotides, deoxyribonucleotides and modified nucleotides such as inosine or nucleotides containing modified groups, which do not essentially alter their hybridization characteristics.

Probe sequences are represented throughout the specification as single stranded DNA oligonucleotides from the 5' to the 3' end. It is obvious to the man skilled in the art that any of the below-specified probes can be used as such, or in their complementary form, or in their RNA form (wherein U replaces T).

The probes according to the invention can be prepared by cloning of recombinant plasmids containing inserts including the corresponding nucleotide sequences, if need be by cleaving the latter out from the cloned plasmids upon using the adequate nucleases and recovering them, e.g. by fractionation according to molecular weight. The probes according to the present invention can also be synthesized chemically, for instance by the conventional phospho-triester method.

The term "solid support" can refer to any substrate to which an oligonucleotide probe can be coupled, provided that it retains its hybridization characteristics and provided that the background level of hybridization remains low. Usually the solid substrate will be a microtiter plate, a membrane (e.g. nylon or nitrocellulose) or a microsphere (bead) or a chip. Prior to application to the membrane or fixation it may be convenient to modify the nucleic acid probe in order to facilitate fixation or improve the hybridization efficiency. Such modifications may encompass homopolymer tailing, coupling with different reactive groups such as aliphatic groups, NH₂ groups, SH groups, carboxylic groups, or coupling with biotin, haptens or proteins.

The term "labeled" refers to the use of labeled nucleic acids. Labeling may be carried out by the use of labeled nucleotides incorporated during the polymerase step of the amplification such as illustrated by Saiki et al. (1988) or Bej et al. (1990) or labeled primers, or by any other method known to the person skilled in the art. The nature of the label may be isotopic (32P, 35S, etc.) or non-isotopic (biotin, digoxigenin, etc.).

The term "primer" refers to a single stranded oligonucleotide sequence capable of acting as a point of initiation for synthesis of a primer extension product, which is complementary to the nucleic acid strand to be copied. The length and the sequence of the primer must be such that they allow to prime the synthesis of the extension products. Preferably the primer is about 5-50 nucleotides long. Specific length and sequence will depend on the complexity of the required DNA or RNA targets, as well as on the conditions of primer use such as temperature and ionic strength.

The term "primer pair" refers to a set of primers comprising at least one 5' primer and one 3' primer. The primer pair may consist of more than two primers, the complexity of the number of primers will depend on the hybridization conditions, variability of the sequences in the regions to be amplified and the target sequences to be detected.

The fact that amplification primers do not have to match exactly with the corresponding template

10

. 15

20

25

30

35

sequence to warrant proper amplification is amply documented in the literature (Kwok et al., 1990).

The amplification method used can be either polymerase chain reaction (PCR; Saiki et al., 1988), ligase chain reaction (LCR; Landgren et al., 1988; Wu & Wallace, 1989; Barany, 1991), nucleic acid sequence-based amplification (NASBA; Guatelli et al., 1990; Compton, 1991), transcription-based amplification system (TAS; Kwoh et al., 1989), strand displacement amplification (SDA; Duck, 1990) or amplification by means of QB replicase (Lomeli et al., 1989) or any other suitable method to amplify nucleic acid molecules known in the art.

The oligonucleotides used as primers or probes may also comprise nucleotide analogues such as phosphorothiates (Matsukura et al., 1987), alkylphosphorothiates (Miller et al., 1979) or peptide nucleic acids (Nielsen et al., 1991; Nielsen et al., 1993) or may contain intercalating agents (Asseline et al., 1984).

As most other variations or modifications introduced into the original DNA sequences of the invention these variations will necessitate adaptations with respect to the conditions under which the oligonucleotide should be used to obtain the required specificity and sensitivity. However the eventual results of hybridization will be essentially the same as those obtained with the unmodified oligonucleotides.

The introduction of these modifications may be advantageous in order to positively influence characteristics such as hybridization kinetics, reversibility of the hybrid-formation, biological stability of the oligonucleotide molecules, etc.

The "sample" may be any biological material taken either directly from the infected human being (or animal), or after culturing (enrichment). Biological material may be e.g. expectorations of any kind, broncheolavages, blood, skin tissue, biopsies, sperm, lymphocyte blood culture material, colonies, liquid cultures, fecal samples, urine etc.

The sets of probes of the present invention will include at least 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30 or more probes. Said probes may be applied in two or more distinct and known positions on a solid substrate. Often it is preferable to apply two or more probes together in one and the same position of said solid support.

For designing probes with desired characteristics, the following useful guidelines known to the person skilled in the art can be applied.

Because the extent and specificity of hybridization reactions such as those described herein are affected by a number of factors, manipulation of one or more of those factors will determine the exact sensitivity and specificity of a particular probe, whether perfectly complementary to its target or not. The importance and effect of various assay conditions, explained further herein, are known to those skilled in the art.

The stability of the [probe: target] nucleic acid hybrid should be chosen to be compatible with the assay conditions. This may be accomplished by avoiding long AT-rich sequences, by terminating the

10

15

20

25

30

35

hybrids with G:C base pairs, and by designing the probe with an appropriate Tm. The beginning and end points of the probe should be chosen so that the length and %GC result in a Tm about 2-10°C higher than the temperature at which the final assay will be performed. The base composition of the probe is significant because G-C base pairs exhibit greater thermal stability as compared to A-T base pairs due to additional hydrogen bonding. Thus, hybridization involving complementary nucleic acids of higher G-C content will be stable at higher temperatures.

Conditions such as ionic strength and incubation temperature under which a probe will be used should also be taken into account when designing a probe. It is known that hybridization will increase as the ionic strength of the reaction mixture increases, and that the thermal stability of the hybrids will increase with increasing ionic strength. On the other hand, chemical reagents, such as formamide, urea, DMSO and alcohols, which disrupt hydrogen bonds, will increase the stringency of hybridization. Destabilization of the hydrogen bonds by such reagents can greatly reduce the Tm. In general, optimal hybridization for synthetic oligonucleotide probes of about 10-50 bases in length occurs approximately 5°C below the melting temperature for a given duplex. Incubation at temperatures below the optimum may allow mismatched base sequences to hybridize and can therefore result in reduced specificity.

It is desirable to have probes, which hybridize only under conditions of high stringency. Under high stringency conditions only highly complementary nucleic acid hybrids will form; hybrids without a sufficient degree of complementarity will not form. Accordingly, the stringency of the assay conditions determines the amount of complementarity needed between two nucleic acid strands forming a hybrid. The degree of stringency is chosen such as to maximize the difference in stability between the hybrid formed with the target and the nontarget nucleic acid. In the present case, single base pair changes need to be detected, which requires conditions of very high stringency.

The length of the target nucleic acid sequence and, accordingly, the length of the probe sequence can also be important. In some cases, there may be several sequences from a particular region, varying in location and length, which will yield probes with the desired hybridization characteristics. In other cases, one sequence may be significantly better than another that differs merely by a single base. While it is possible for nucleic acids that are not perfectly complementary to hybridize, the longest stretch of perfectly complementary base sequence will normally primarily determine hybrid stability. While oligonucleotide probes of different lengths and base composition may be used, preferred oligonucleotide probes of this invention are between about 5 to 50 (more particularly 10-25) bases in length and have a sufficient stretch in the sequence which is perfectly complementary to the target nucleic acid sequence.

Regions in the target DNA or RNA, which are known to form strong internal structures inhibitory to hybridization, are less preferred. Likewise, probes with extensive self-complementarity should be avoided. As explained above, hybridization is the association of two single strands of complementary nucleic acids to form a hydrogen bonded double strand. It is implicit that if one of the two strands is wholly or partially involved in a hybrid that it will be less able to participate in formation

10

15

20

30

35

of a new hybrid. There can be intramolecular and intermolecular hybrids formed within the molecules of one type of probe if there is sufficient self complementarity. Such structures can be avoided through careful probe design. By designing a probe so that a substantial portion of the sequence of interest is single stranded, the rate and extent of hybridization may be greatly increased. Computer programs are available to search for this type of interaction. However, in certain instances, it may not be possible to avoid this type of interaction.

Standard hybridization and wash conditions are disclosed in the Materials & Methods section of the Examples. Other conditions are for instance 3X SSC (Sodium Saline Citrate), 20% deionized FA (Formamide) at 50°C.

Other solutions (SSPE (Sodium saline phosphate EDTA), TMACI (Tetramethyl ammonium Chloride), etc.) and temperatures can also be used provided that the specificity and sensitivity of the probes is maintained. If need be, slight modifications of the probes in length or in sequence have to be carried out to maintain the specificity and sensitivity required under the given circumstances.

Primers may be labeled with a label of choice (e.g. biotin). Different primer-based target amplification systems may be used, and preferably PCR-amplification, as set out in the examples. Single-round or nested PCR may be used.

The term "hybridization buffer" means a buffer enabling a hybridization reaction to occur between the probes and the polynucleic acids present in the sample, or the amplified products, under the appropriate stringency conditions.

The term "wash solution" means a solution enabling washing of the hybrids formed under the appropriate stringency conditions.

The following examples only serve to illustrate the present invention. These examples are in no way intended to limit the scope of the present invention.

25 FIGURE AND TABLE LEGENDS

Figure 1: Natural and drug selected variability in the vicinity of codons 30, 46, 48, 50, 54, 82, 84, and 90 of the HIV-1 protease gene. The most frequently observed wild-type sequence is shown in the top line. Naturally occurring variations are indicated below and occur independently from each other.

Drug-selected variants are indicated in bold

Figure 2 A: Reactivities of the selected probes for codon 30 immobilized on LiPA strips with reference material. The information in the boxed surface is not relevant for the discussion of probes for condon 30 The position of each selected probe on the membrane strip is shown at the left of each panel. The sequence of the relevant part of the selected probes is shown at the left and is given in Table 1. Each strip is incubated with a biotinylated PCR fragment from the reference panel. The reference panel accession

20

35

numbers are indicated in <u>Table 1</u>. For several probes multiple reference panel possibilities are available, but only one relevant accession number given each time. *: False positive reactivities. At the bottom the strips, the amino acids at the relevant codon, as derived from the probe reactivity, is indicated.

- Figure 2 B: Reactivities of the selected probes for codons 46 and 48 immobilized on LiPA strips with reference material. The information in the boxed surface is not relevant for the discussion of probes for condons 46 and 48 The position of each selected probe on the membrane strip is shown at the left of each panel. The sequence of the relevant part of the selected probes is given in Table 1. Each strip is incubated with a biotinylated PCR fragment from the reference panel. The reference panel accession numbers are indicated in Table 1. For several probes multiple reference panel possibilities are available, but only one relevant accession number given each time. *: False positive reactivities. On top of the strips, the amino acids at the relevant codon, as derived from the probe reactivity, is indicated.
 - Figure 2 C: Reactivities of the selected probes for codon 50 immobilized on LiPA strips with reference material. The information in the boxed surface is not relevant for the discussion of probes for condon 50. The position of each selected probe on the membrane strip is shown at the left of each panel. The sequence of the relevant part of the selected probes is given in Table 1. Each strip is incubated with a biotinylated PCR fragment from the reference panel. The reference panel accession numbers are indicated in <u>Table 1</u>. For several probes multiple reference panel possibilities are available, but only one relevant accession number given each time. *: False positive reactivities. At the bottom of the strips, the amino acids at the relevant codon, as derived from the probe reactivity, is indicated.
- Figure 2 D: Reactivities of the selected probes for codon 54 immobilized on LiPA strips with reference material. The information in the boxed surface is not relevant for the discussion of probes for condon 54.

 The position of each selected probe on the membrane strip is shown at the left of each panel. The sequence of the relevant part of the selected probes is given in Table 1. Each strip is incubated with a biotinylated PCR fragment from the reference panel. The reference panel accession numbers are indicated in Table 1. For several probes multiple reference panel possibilities are available, but only one relevant accession number given each time. *: False positive reactivities. At the bottom of the strips, the amino acids at the relevant codon, as derived from the probe reactivity, is indicated.
 - Figure 2 E.:Reactivities of the selected probes for codons 82 and 84 immobilized on LiPA strips with reference material. The information in the boxed surface is not relevant for the discussion of probes for condons 82 and 84. The position of each selected probe on the membrane strip is shown at the left of each panel. The sequence of the relevant part of the selected probes is given in Table 1. Each strip is incubated with a biotinylated PCR fragment from the reference panel. The reference panel accession

25

30

35

numbers are indicated in <u>Table 1</u>. For several probes multiple reference panel possibilities are available, but only one relevant accession number given each time. *: False positive reactivities. At the bottom of the strips, the amino acids at the relevant codon, as derived from the probe reactivity, is indicated.

- Figure 2 F: Reactivities of the selected probes for codon 90 immobilized on LiPA strips with reference material. The information in the boxed surface is not relevant for the discussion of probes for condon 90. The position of each selected probe on the membrane strip is shown at the left of each panel. The sequence of the relevant part of the selected probes is given in Table 1. Each strip is incubated with a biotinylated PCR fragment from the reference panel. The reference panel accession numbers are indicated in Table 1. For several probes multiple reference panel possibilities are available, but only one relevant accession number given each time. *: False positive reactivities. At the bottom of the strips, the amino acids at the relevant codon, as derived from the probe reactivity, is indicated.
 - Figure 3: Sequence and position of the HIV-1 protease amplification primers. To obtain the reactivity with probes selected to determine the susceptibility to antiviral drugs involving codons 30, 46, 48, 50, 54, 82, and 84, nested amplification primers prot2bio(5' primer) and Prot31bio (3' primer) were designed. To obtain the reactivity with probes selected to determine the susceptibility to antiviral drugs involving codon 90, nested amplification primers Prot41bio (5' primer) and Prot6bio (3' primer) were designed.
- Figure 4 A: Phylogenetic analysis on 312 protease sequences allowed to separate genotype B strains from non-B strains. Reactivities of the selected probes for codon 30 immobilized on LiPA strips with a biotinylated PCR fragment of genotype B strains and non-B strains is shown, the exact percentages are indicated in table 5. The probes are indicated at the bottom. The sequence of the relevant part of the probes is given in Table 1.

Figure 4 B: Phylogenetic analysis on 312 protease sequences allowed to separate genotype B strains from non-B strains. Reactivities of the selected probes for codons 46/48 immobilized on LiPA strips with a biotinylated PCR fragment of genotype B strains and non-B strains is shown, the exact percentages are indicated in table 5. The probes are indicated at the bottom. The sequence of the relevant part of the probes is given in Table 1.

Figure 4 C: Phylogenetic analysis on 312 protease sequences allowed to separate genotype B strains from non-B strains. Reactivities of the selected probes for codon 50 immobilized on LiPA strips with a biotinylated PCR fragment of genotype B strains and non-B strains is shown, the exact percentages are indicated in table 5. The probes are indicated at the bottom. The sequence of the relevant part of the probes is given in Table 1.

10

25

Figure 4 D: Phylogenetic analysis on 312 protease sequences allowed to separate genotype B strains from non-B strains. Reactivities of the selected probes for codon 54 immobilized on LiPA strips with a biotinylated PCR fragment of genotype B strains and non-B strains is shown, the exact percentages are indicated in table 5. The probes are indicated at the bottom. The sequence of the relevant part of the probes is given in Table 1.

Figure 4 E: Phylogenetic analysis on 312 protease sequences allowed to separate genotype B strains from non-B strains. Reactivities of the selected probes for codons 82/84 immobilized on LiPA strips with a biotinylated PCR fragment of genotype B strains and non-B strains is shown, the exact percentages are indicated in table 5. The probes are indicated at the bottom. The sequence of the relevant part of the probes is given in Table 1.

- Figure 4 F: Phylogenetic analysis on 312 protease sequences allowed to separate genotype B strains from non-B strains. Reactivities of the selected probes for codon 90 immobilized on LiPA strips with a biotinylated PCR fragment of genotype B strains and non-B strains is shown, the exact percentages are indicated in table 5. The probes are indicated at the bottom. The sequence of the relevant part of the probes is given in Table 1.
- Figure 5 A: Geographical origin of 856 samples and reactivities with the different probes at codon position 30. The exact percentages are indicated in table 6. The probes are indicated at the bottom.
 - Figure 5 B: Geographical origin of 856 samples and reactivities with the different probes at codon positions 46/48. The exact percentages are indicated in table 6. The probes are indicated at the bottom.
 - Figure 5 C: Geographical origin of 856 samples and reactivities with the different probes at codon position 50. The exact percentages are indicated in table 6. The probes are indicated at the bottom.
- Figure 5 D: Geographical origin of 856 samples and reactivities with the different probes at codon position 54. The exact percentages are indicated in table 6. The probes are indicated at the bottom.
 - Figure 5 E: Geographical origin of 856 samples and reactivities with the different probes at codon positions 82/84. The exact percentages are indicated in table 6. The probes are indicated at the bottom.
- Figure 5 F: Geographical origin of 856 samples and reactivities with the different probes at codon position 90. The exact percentages are indicated in table 6. The probes are indicated at the bottom.

Table 1: HIV-1 protease wild-type and drug-selected mutation probes with their corresponding sequences as applied on the HIV-1 protease LiPA strip. The most frequently observed wild-type sequence is shown at the top line. Probe names corresponding to the selected motifs are indicated in the left column, the relevant part of each probe applied on the strip is shown under the consensus sequence.

Table 2: Protease Inhibitors.

5

Table 3: HIV-1 protease wild-type and drug-selected mutation probes with their corresponding sequences as synthesized, immobilized and tested on LiPA strips. The most frequently observed wild-type sequence is shown at the top line. Probe names corresponding to the selected motifs are indicated in the left column, the relevant part of each probe applied on the strip is shown under the consensus sequence. The probes retained are indicated in table 1.

15 Table 4: Polymorphic nucleotide sequences.

Table 5: % Reactivities of the HIV-1 protease wild-type and drug-selected mutation probes applied on the HIV-1 protease LiPA strip with genotype B strains and non-B strains.

Table 6: % Reactivities of the HIV-1 protease wild-type and drug-selected mutation probes applied on the HIV-1 protease LiPA strip with samples of different geographical origin.

Table 7: HIV-1 protease wild-type and drug-selected mutation probes with their corresponding sequences as applied on the HIV-1 protease LiPA strip. The most frequently observed wild-type sequence is shown at the top line. Probe names corresponding to the selected motifs are indicated in the left column, the relevant part of each probe applied on the strip is shown under the consensus sequence.

EXAMPLES

30 Example 1:

25

35

Selection of the plasma samples, PCR amplification and cloning of the PCR products.

<u>Plasma samples</u> (n=557) were taken from HIV type-1 infected patients and stored at -20°C until use. Plasma samples were obtained from naive and drug-treated patients. The drugs involved ritonavir, indinavir and saquinavir. The serum samples were collected from patients residing in Europe (Belgium, Luxembourg, France, Spain and UK), USA and Brazil.

WO 99/67428 PCT/EP99/04317

HIV RNA was prepared from these samples using the guanidinium-phenol procedure. Fifty µl plasma was mixed with 150 µl Trizol®LS Reagent (Life Technologies, Gent, Belgium) at room temperature (volume ratio: 1unit sample/ 3 units Trizol). Lysis and denaturation occurred by carefully pipetting up and down several times, followed by an incubation step at room temperature for at least 5 minutes. Fourthy µl CHCl₃ was added and the mixture was shaken vigorously by hand for at least 15 seconds, and incubated for 15 minutes at room temperature. The samples were centrifuged at maximum 12,000g for 15 minutes at 4°C, and the colorless aqueous phase was collected and mixed with 100 µl isopropanol. To visualize the minute amounts of viral RNA, 20 µl of 1µg/µl Dextran T500 (Pharmacia) was added, mixed and left at room temperature for 10 minutes. Following centrifugation at max. 12,000g for 10 minutes at 4°C and aspiration of the supernatant, the RNA pellet was washed with 200 µl ethanol, mixed by vortexing and collected by centrifugation at 7,500g for 5 minutes at 4°C. Finally the RNA pellet was briefly air-dried and stored at -20°C. Alternatively, the High Pure Viral Nucleic Acid Kit (Boehringer Mannheim) was used to extract RNA from the samples

5

10

15

20

25

30

35

For cDNA synthesis and PCR amplification, the RNA pellet was dissolved in 15 μl random primers (20 ng/μl, pdN₆, Pharmacia), prepared in DEPC-treated or HPLC grade water. After denaturation at 70°C for 10 minutes, 5 μl cDNA mix was added, composed of 4 μl 5x AMV-RT buffer (250mM Tris.HCl pH 8.5, 100mM KCl, 30mM MgCl₂, 25 mM DTT), 0.4 μL 25mM dXTPs, 0.2 μl or 25U Ribonuclease Inhibitor (HPRI, Amersham), and 0.3 μl or 8U AMV-RT (Stratagene). cDNA synthesis occurred during the 90 minutes incubation at 42°C. The HIV -1 protease gene was than amplified using the following reaction mixture: 5 μl cDNA, 4.5 μl 10x Taq buffer, 0.3 μl 25 mM dXTPs, 1 μl (10 pmol) of each PCR primer, 38 μl H₂O, and 0.2 μl (1 U) Taq. . Alternatively, the Titon One Tube RT-PCR system (Boehringer Mannheim) was used to perform RT-PCR.

Codon positions involving resistance to saquinavir, ritonavir, indinavir, nelfinavir and VX-478 have been described (Shinazi et al) and PCR amplification primers were chosen outside these regions. The primer design was based on HIV-1 published sequences (mainly genotype B clade) (Myers et al.) and located in regions that showed a high degree of nucleotide conservation between the different HIV-1 clades. The final amplified region covered the HIV-1 protease gene from codon 9 to codon 99. The primers for the following sequence: outer sense primer CAGAGCCAACAGCCCCACCAG3' (SEQ ID NO 1); nested sense primer Prot 2 bio: 5' CCT CAR ATC ACT CTT TGG CAA CG 3' (SEQ ID NO 3); nested antisense primer Prot 6 bio: 3' TAA TCR GGA TAA CTY TGA CAT GGT C 5' (SEQ ID NO 4); and outer antisense primer RT12: 5' bioATCAGGATGGAGTTCATAACCCATCCA3' (SEQ ID NO 2). Annealing occurred at 57°C, extension at 72°C and denaturation at 94°C. Each step of the cycle took 1 minute, the outer PCR contained 40 cycles, the nested round 35. Nested round PCR products were analyzed on agarose gel and only clearly visible amplification products were used in the LiPA procedure. Quantification of viral

10

15

20

25

35

RNA was obtained with the HIV MonitorTMtest (Roche, Brussels, Belgium). Later on, new sets of primers for amplification were selected. For the amplification of HIV protease codon 30-84: outer sense primer prot16: 5'-CAGAGCCAACAGCCCCACCAG-3' (SEQ ID NO 501), outer antisense primer prot5: 5'-TTTTCTTCTGTCAATGGCCATTGTTT-3' (SEQ ID NO 502) were used. Annealing occurred at 50°C, extension at 68°C and denaturation at 94°C for 35 cycles for the outer PCR. For the nested PCR annealing occurred at 45°C, denaturation at 94°C and extension at 92°C with primers: nested sense primers prot2a-bio: 5'-bio-CCTCAAATCACTCTTTGGCAACG-3' (SEQ ID NO 503)and prot2b-bio: 5'-bio-CCTCAGATCACTCTTTGGCAACG-3' (SEQ ID NO 504), and nested antisense primer prot31bio: 5'-bio-AGTCAACAGATTTCTTCCAAT-3' (SEQ ID NO 6). For the amplification of HIV protease codon 90, the outer PCR was as specified for HIV protease codon 30-84. For the nested PCR, nested sense primer prot41-bio: 5'-bio-CCTGTCAACATAATTGCAAG-3' (SEQ ID NO 505) and nested antisense primers prot6a: 5'-bio-CTGGTACAGTTTCAATAGGGCTAAT-3' (SEQ ID NO 506), prot6b: (SEQ ID NO prot6c: 5'-bio-507), 5'-bio-CTGGTACAGTTTCAATAGGACTAAT-3' CTGGTACAGTCTCAATAGGACTAAT-3' IDNO 508), prot6d: 5'-bio-(SEQ CTGGTACAGTCTCAATAGGGCTAAT-3' (SEQ ID NO 509) were used. For the nested PCR the annealing temperature occurred at 45°C. Primers were tested on a plasmid, which contained an HIV fragment of 1301 bp ligated in a pGEM-T vector. The fragment contains protease, reverse transcriptase and the primer sites of first and second round PCR. By restriction with Sac I the plasmid is linearised.

Selected PCR products were cloned into the pretreated EcoRV site of the pGEMT vector (Promega). Recombinant clones were selected after α-complementation and restriction fragment length analysis, and sequenced using standard sequencing techniques with plasmid primers and internal HIV protease primers. Sometimes biotinylated fragments were directly sequenced with a dye-terminator protocol (Applied Biosystems) using the amplification primers. Alternatively, nested PCR was carried out with analogs of the nested primers, in which the biotin group was replaced with the T7- and SP6-primer sequence, respectively. These amplicons were than sequenced with an SP6- and T7-dye-primer procedure.

Example 2:

30 Selection of a reference panel

Codon positions involving resistance to saquinavir, ritonavir, indinavir, nelfinavir and VX-478 have been described (Shinazi et al. 1997). It was the aim to clone in plasmids those viral protease genes that are covering the different genetic motifs at those important codon positions conferring resistance against the described protease inhibitors.

After careful analysis of 312 protease gene sequences, obtained after direct sequencing of PCR fragments, a selection of 47 PCR fragments which covered the different target polymorphisms and

10

15

20

25

30

35

mutations were retained and cloned in plasmids using described cloning techniques. The selection of samples originated from naive or drug-treated European, Brazilian or US patients. These 47 recombinant plasmids are used as a reference panel, a panel that was sequenced on both strands, and biotinylated PCR products from this panel were used to optimize probes for specificity and sensitivity.

Although this panel of 47 samples is a representative selection of clones at this moment, it is important to mention here that this selection is an fact only a temporally picture of the variability of the virus, and a continuous update of this panel will be mandatory. This includes on ongoing screening for the new variants of the virus, and recombinant cloning of these new motifs.

Probe selection and LiPA testing.

To cover all the different genetic motifs in the reference panel, a total of 471 probes were designed (codon 30: 40 probes; codon 46/48: 72 probes; codon 50:55 probes; codon 54: 54 probes; codon 82/84: 130 probes; codon 90: 120 probes). Table 3 shows the different probes that were selected for the different codon positions.

It was the aim to adapt all probes to react specifically under the same hybridization and wash conditions by carefully considering the % (G+C), the probe length, the final concentration of the buffer components, and hybridization temperature (Stuyver et al., 1997). Therefore, probes were provided enzymatically with a poly-T-tail using the TdT (Pharmacia) in a standard reaction condition, and purified via precipitation. For a limited number of probes with 3' T-ending sequences, an additional G was incorporated between the probe sequence and the poly-T-tail in order to limit the hybridizing part to the specific probe sequence and to exclude hybridization with the tail sequence. Probe pellets were dissolved in standard saline citrate (SSC) buffer and applied as horizontal parallel lines on a membrane strip. Control lines for amplification (probe 5' TAGGGGGAATTGGAGGTTTTAG 3', HIV protease aa 47 to aa 54) and conjugate incubation (biotinylated DNA) were applied alongside. Probes were immobilized onto membranes by baking, and the membranes were sliced into 4mm strips also called LiPA strips.

Selection of the amplification primers and PCR amplification was as described in example 1. In order to select specific reacting probes out of the 471 candidate probes, LiPA tests were performed with biotinylated PCR fragments from the reference panel. To perform LiPA tests, equal amounts (10 µl) of biotinylated amplification products and denaturation mixture (0.4 N NaOH/0.1% SDS) were mixed, followed by an incubation at room temperature for 5 minutes. Following this denaturation step, 2 ml hybridization buffer (2xSSC, 0.1% SDS, 50mM Tris pH7.5) was added together with a membrane strip and hybridization was carried out at 39°C for 30 min. Then, the hybridization mixture was replaced by stringent washing buffer (same composition as hybridization buffer), and stringent washing occurred first at room temperature for 5 minutes and than at 39°C for another 25 minutes. Buffers were than replaced to be suitable for the streptavidine alkaline phosphatase conjugate incubations. After 30 minutes

incubation at room temperature, conjugate was rinsed away and replaced by the substrate components for alkaline phosphatase, Nitro-Blue-Tetrazolium and 5-Bromo-4-Chloro-3-Indolyl Phosphate. After 30 minutes incubation at room temperature, probes where hybridization occurred became visible because of the purple brown precipitate at these positions.

After careful analysis of the 471 probes, the most specific and sensitive probes (n=46) were finally selected, covering the natural and drug-selected variability in the vicinity of aa. 30, 46, 48, 50, 54, 82, 84, and 90. Figure 2 shows the reactivity of the finally selected probes with the reference panel.

Example 3:

5

15

20

25

30

10 LiPA testing on clinical samples.

A total of 856 samples were tested on this selection of 46 specific probes. The geographical origin of these samples is as follows: USA:359; France: 154; UK:36; Brazil 58; Spain 35; Belgium 199; Luxembourg: 15.

From this population, a total of 144 samples were sequenced which allowed to separate the genotype B samples (94) from the non-B samples (50). After analysis of these genotyped samples on LiPA, the genotypic reactivity on the selected probes was scored. Figures 4A to 4F show these results for the different codon positions and for the genotype B versus non-B group. From these tables, it is clear that there is little difference in sequence usage for the different codon positions with respect to specific reactivities at the different probes.

The total collection of 856 samples was then tested on the available 46 probes. After dissection of these reactivities over the different probes and different geographical origin, the picture looks as is presented in Figures 5A to 5F. Again here, the majority of the sequences used at the different codon positions are restricted to some very abundant wild type motifs. It is important to mention here that the majority of these samples are taken from patients never treated with protease inhibitors, en therefore, the majority of the reactivities are found in wild type motifs. Nevertheless, it is clear from some codon positions that the variability at some codon positions in the mutant motif might be considerable, and again, a continuos update on heavily treated patients is mandatory. Another issue is the amount of double blank reactivities, which is in this approach reaching up to 5% in global; with some peak values for some countries for some codon positions: for example 13.8% for codon 82/85 in Brazil; and 18.1 % for codon 90 in Belgium.

The continuous update resulted in a further selection of probes. This later configuration of the strip is indicated in table 7.

Asseline U, Delarue M, Lancelot G, Toulme F, Thuong N (1984) Nucleic acid-binding molecules with high affinity and base sequence specificity: intercalating agents covalently linked to oligodeoxynucleotides. Proc. Natl. Acad. Sci. USA 81(11):3297-301.

Barany F. Genetic disease detection and DNA amplification using cloned thermostable ligase. Proc Natl Acad Sci USA 1991; 88: 189-193.

Bej A, Mahbubani M, Miller R, Di Cesare J, Haff L, Atlas R. Mutiplex PCR amplification and immobilized capture probes for detection of bacterial pathogens and indicators in water. Mol Cell Probes 1990; 4:353-365.

10

Compton J. Nucleic acid sequence-based amplification. Nature 1991; 350: 91-92.

Condra et al. 1995. In vivo emergence of HIV-1 variants resistant to multiple protease inhibitors. Nature 374: 569-571.

15

Condra et al. 1995. In vivo emergence of HIV-1 variants resistant to multiple protease inhibitors. Nature 374: 569-571.

Duck P. Probe amplifier system based on chimeric cycling oligonucleotides. Biotechniques 1990; 9: 142-20 147.

Eberle et al. 1995. Resistance of HIV type 1 to proteinase inhibitor Ro 31-8959. AIDS Research and Human Retroviruses 11: 671-676.

- Emini et al. 1994. Phenotypic and genotypic characterization of HIV-1 variants selected during treatment with the protease inhibitor L-735, L-524. Third International Workshop on HIV Drug Resistance, Kauai, Hawaii, USA.
- Guatelli J, Whitfield K, Kwoh D, Barringer K, Richman D, Gengeras T. Isothermal, in vitro amplification of nucleic acids by a multienzyme reaction modeled after retroviral replication. Proc Natl Acad Sci USA 1990; 87: 1874-1878.

Huff et al. 1991. HIV protease: a novel chemotherapeutic target for AIDS. J. Med. Chem. 34: 2305-2314.

35

Hunter et al. 1994. Macromolecular interactions in the assembly of HIV and other retroviruses.

25

Seminars in Virology 5: 71-83.

Kempf et al. 1994. Pharmacokinetic and in vitro selection studies with ABT-538, a potent inhibitor of HIV protease with high oral bioavailability. 34th Interscience Conference on Antimicrobial Agents and Chemotherapy, Orlando, Fl, USA.

Kohl et al. 1988. Active human immunodeficiency virus protease is required for viral infectivity. Proc. Natl. Acad. Sci. USA. 85: 4686-4690.

- 10 Kwok S, Kellogg D, McKinney N, Spasic D, Goda L, Levenson C, Sinisky J. Effects of primer-template mismatches on the polymerase chain reaction: Human immunodeficiency views type 1 model studies. Nucl. Acids Res. 1990; 18: 999.
- Landgren U, Kaiser R, Sanders J, Hood L. A ligase-mediated gene detection technique. Science 1988; 241:1077-1080.
 - Lomeli H, Tyagi S, Printchard C, Lisardi P, Kramer F. Quantitative assays based on the use of replicatable hybridization probes. Clin Chem 1989; 35: 1826-1831.
- 20 Matsukura M, Shinozuka K, Zon G, Mitsuya H, Reitz M, Cohen J, Broder S (1987) Phosphorothioate analogs of oligodeoxynucleotides: inhibitors of replication and cytopathic effects of human immunodeficiency virus. Proc. Natl. Acad. Sci. USA 84(21):7706-10.
 - Meek et al. 1989. Proc. Natl. Acad. Sci. USA. 86: 1841-1845.
- Miller P, Yano J, Yano E, Carroll C, Jayaram K, Ts'o P (1979) Nonionic nucleic acid analogues. Synthesis and characterization of dideoxyribonucleoside methylphosphonates. Biochemistry 18(23):5134-43.
- Myers et al. 1996. Human retroviruses and AIDS 1996. Los Alamos Laboratory, Los Alamos, N.M. Navia et al. 1989. Three-dimensional structure of aspartyl protease from human immunodeficiency virus HIV-1. Nature 337: 615-620.
- Nielsen P, Egholm M, Berg R, Buchardt O (1993) Sequence specific inhibition of DNA restriction enzyme cleavage by PNA. Nucleic-Acids-Res. 21(2):197-200.

Nielsen P, Egholm M, Berg R, Buchardt O (1991) Sequence-selective recognition of DNA by strand displacement with a thymine-substituted polyamide. Science 254(5037):1497-500.

Patick et al. 1996. Antiviral and resistance studies of AG1343, an orally bioavailable inhibitor of human immunodeficiency virus protease. Antimicrobial Agents and Chemotherapy 40: 292-297; 40: 1575 (erratum).

Peng et al. 1989. Role of human immunodeficiency virus type 1-specific protease in core protein maturation and viral infectivity. J. Virol. 63: 2550-2556.

10

Rao et al. 1996. Structural and modeling analysis of the basis of viral resistance to VX-478. Fifth International Workshop on HIV Drug Resistance, Whistler, Canada, abstract n°: 22.

Saiki R, Walsh P, Levenson C, Erlich H. Genetic analysis of amplified DNA with immobilized sequencespecific oligonucleotide probes Proc Natl Acad Sci USA 1989; 86:6230-6234.

Schmit et al. 1996. Resistance-related mutations in the HIV-1 protease gene of patients treated for 1 year with the protease inhibitor ritonavir (ABT-538). AIDS 10: 995-999.

20 Shinazi et al. 1997. Mutations in retroviral genes associated with drug resistance. International Antiviral News 5: 129-142.

Stuyver L, Rossau R, Wyseur A, et al. Typing of hepatitis C virus isolates and characterization of new subtypes using a line probe assay. J. Gen. Virol. 1993; 74: 1093-1102.

25

Tisdale et al. 1994. Comprehensive analysis of HIV-1 variants individually selected for resistance to six HIV protease inhibitors. Third International Workshop on HIV Drug Resistance, Kauai, Hawaii, USA.

30

Wlodawer et al. 1993. Structure-based inhibitors of HIV-1 protease. Annu. Rev. Biochem. 62:543-585.

Wlodawer et al. 1989. Science 245: 616-621.

Wu D, Wallace B. The ligation amplification reaction (LAR) - amplification of specific DNA sequences using sequential rounds of template-dependent ligation. Genomics 1989; 4:560-569.

Table 1

	26 27 ACA GGA	28 29 GCA GAT	30 31 GAT ACA	32 GTA	33 TTA	34 GAA		lengte	Seq ID
pc30w25 pc30w29 pc30w32 pc30w36 pc30m23	. А	GCA GAT	GAT ACA GAT ACA GAC ACA GAT ACA	GT GT GG			40 36 42 40 40	14 13 14 14 15	31 35 38 42 29
pc48w47 pc48w45 pc48w72 pc48m41	AAA A	46 47 ATG ATA ATG ATA ATG ATA ATA ATA ATG ATA	GGG GGA GGG GGA	ATT	51 GGA		42 42 42 40	15 16 16 15	93 91 120 87
pc50w31 pc50w44 pc50w52 pc50m37	GGA A GGA A GA A	ATT GGA G ATT GGA G ATT GGG G ATT GGA G	GT TTT	54 ATC			42 42 40	15 15 14 12	151 164 172 157
pc54w3 pc54w34 pc54w14 pc54w19 pc54w22 pc54w26 pc54w27 pc54w35 pc54m35	GT GA GGT GA GGT GA GGT GGT GGT GGT GGT	ITT ATT A ITC ATT A ITT ATT A ITT GCC A ITT GTC A	AAA GTA	AGA A	58 CAG		42 42 42 42 42 42 40 38 40 42	17 16 16 17 16 15 15	178 212 189 194 197 202 204 213 215
pc82w91 pc82w60 pc82w111 pc82w89 pc82w42 pc82m36 pc82m67 pc82m38 pc82m10 pc82m10 pc82m63 pc82m10	1 5 7	A CCG ACA CCT ACA CCT ACA CCT ACA CCT ACA CCT ACG CCC CA CCT ACA CCT	GTC AAC GTC AAC GTT AAC GTC AAC ACC AAC ACC AAC TTC AAC TTC AAC TTC AAC GCC AAC	ATA ATA ATA GTA ATA ATA ATA ATA ATA	A ATG ATT AG	86 GGA	87 AGA 44 42 42 42 42 40 44 44 44	16 17 16 17 14 15 15 15 17 15 16 18	318 287 338 316 269 263 294 265 332 354 267 290 328

Table 1 - Cont'd

	86	87	88	89	90	91	92	93	94			
	GGA	AGA	AAT	CTG	TTG	ACT	CAG	ATT	GGT			
pc90w27			AAT	CTG	TTG	ACT	CA			38	3 14	384
pc90w37			AAT	CTG	TTG	ACT	CAG	ATG		42	2 18	394
pc90w39		GA	ACT	CTG	TTG	ACT	С			44	1 15	396
pc90w50			AAT	ATG	TTG	ACT	CAG			. 4(15	407
pc90w52			AAT	TTG	TTG	ACT	CAG			4(15	409
pc90w69		GA	AAC	CTG	TTG	ACT	•			4(14	426
pc90w73				TG	TTG	ACA	CAG	CTT	G	4 4	1 15	430
pc90w79				TG	TTG	ACC	CAG	ATT	G	4.4	15	436
pc90m43		Α	AAT	CTG	ATG	ACT	CA			4 (15	400
pc90m56			AAT	ATG	ATG	ACC	CAG			42	2 15	413

Table 2
Protease Inhibitors

	A mile a said abance	Codon change
Compound	Amino acid change	Codon change
Protease Inhibitors		
A-77003	R8Q	CGA to CAA
	R8K	CGA to AAA
	V32I	GTA to ATA
	M46I	ATG to ATA
	M46L	ATG to TTC
	M46F	ATG to TTC
	M46V	ATG to GTG
	G48V	GGG to GTG
	A71V	GCT to GTT
	V82I	GTC to ATC
·	V82A	GTC to GCC
	L63P	CTC to CCC
	A71T	GCT to ACT
	A71V	GCT to GTT
	G73S	GGT to GCT
	V82A	GTC to GCC
	V82F	GTC to TTC
	V82T	GTC to ACC
	I84V	ATA to GTA
	L90M	TTG to ATG
P9941	V82A	GTC to GCC
Ro 31-8959	L10I	CTC to ATC
(saquinavir)	G48V	GGG to GTG
	I54V	ATC to GTC
	I54V	ATA to GTA
	G73S	GGT to AGT
	V82A	GTC to GCC
	184V	ATA to GTA
	L90M	TTG to ATG
		
RPI-312	I84V	ATA to GTA

29 Table 2 - Cont'd-1

SC-52151	L24V G48V A71V V75I P81T V82A N88D		TTA to GTA GGG to GTG GCT to GTT GTA to ATA CCT to ACT GTC to GCC AAT to GAT
SC-55389A	L10F N88S	-99-	CTC to CGC
SKF108842	V82T I84V		GTC to ACC ATA to GTA
SKF108922	V82A V82T		GTC to GCC GTC to ACC
VB 11,328	L10F M46I I47V I50V 184V		CTC to GGC ATG to ATA ATA to CTA ATT to GTT ATA to GTA
VX-478 (141W94)	L10F M46I I47V I50V I84V		CTC to CGC ATG to ATA ATA to CTA ATT to GTT ATA to GTA
XM323	L10F K45I M46L V82A V82I V82F I84V		CTC to CGC AAA to ATA ATG to CTG GTC to GCC GTC to ATC GTC to TTC ATA to GTA
	L97V 182T		TTA to GTA ATC to ACC
A-75925 ABT-538 (ritonavir)	V32I K20R L33F		GTA to ATA AAG to AAA TTA to TTC

Table 2 - Cont'd-2

	M36I	ATG to ATA
	M46I	ATG to ATA
	154L	ATC to?
	I54V	ATC to GTC
	A71V	GTC to GTT
	V82F	GTC to TTC
	V82A	GTC to GCC
•	V82T	GTC to ACC
	V82S	GTC to TCC
·	I84 V	ATA to GTA
	L90M	TTG to ATG
AG1343		
(nelfinavir)	D30N	GAT to AAT
` '	M36I	•
	M46I	ATG to ATA
	L63P	CTC to CCC
	A71V	GCT to GTT
	V771	
	184 V	ATA to GTA
	N88D	
	L90M	TTG to ATG
BILA 1906	V32I	GTA to ATA
BS	M46I	ATG to ATA
	M46L	ATG to TTG
	A71V	GCT to GTT
	I84A	ATA to GCA
	184V	ATA to GTA
BILA 2011	V32I	GTA to ATA
(palinavir)	A71V	GCT to GTT
	I84A	ATG to ATA
	L63P	CTC to CCC
BILA 2185 BS	L23I	CTA to ATA
BMS 186,318	A71T	GCT to ACT
2112 100,010	V82A	GTC to GCC
DMP 450	L10F	CTC to TTC

Table 2 - Cont'd-3

M46I D60E I84V	ATG to ATA GAT to GAA ATA to GTA
V32I	GTA to ATA
L10I	CTC to ATC
L10R	CTC to CGC
L10V	CTC to GTC
K20M	AAG to ATG
K20R	AAG to AAA
L24I	TTA to ATA
V32I	GTA to ATA
M46I	ATG to ATA
M46L	ATG to TTG
I54V	ATC to GTC
	D60E 184V V32I L10I L10R L10V K20M K20M K20R L24I V32I M46I M46L

Table 3

P30w1 A GCA GAT GAT ACA GTA TT 18 7 P30w2 GA GCA GAT GAT ACA GTA TT 19 8 P30w3 A GCA GAT GAT ACA GTA TT 19 9 P30w4 GGA GCA GAT GAT ACA GTA TT 19 9 P30w5 GGA GCA GAT GAT ACA GTA TT 20 10 P30w6 ACA GGA GCA GAT GAT ACA GTA TTA 21 11 P30w7 CA GGA GCA GAT GAT ACA GTA TTA 18 12 P30w8 A GGA GCA GAT GAT ACA GTA TG 19 13 P30w9 GGA GCA GAT GAT ACA GTA TG 19 15 P30w10 ACA GGA GCA GAT GAT ACA GTA TG 19 16 P30m11 A GCA GAT GAT GAT ACA GTA TT 18 17 P30m12 GA GCA GAT AAT ACA GTA TT 19 18 P30m13 A GCA GAT AAT ACA GTA TT 19 18 P30m14 GGA GCA GAT AAT ACA GTA TT 19 19 P30m15 GCA GAG GCA GAT AAT ACA GTA TT 19 19 P30m16 ACA GGA GCA GAT AAT ACA GTA TT 20 24		26 ACA	27 GGA	28 GCA	29 GAT	30 GAT	31 aca	32 GTA	33 TTT A	34	35	length	Seq ID
P30w2 GA GCA GAT GAT ACA GTA TT 19 8 P30w3 A GCA GAT GAT ACA GTA TTA 19 9 P30w4 GCA GCA GCA GCA GAT ACA GTA TTA 19 9 P30w5 GCA GCA GCA GAT ACA GTA TTA 20 10 P30w6 ACA GGA GCA GAT GAT ACA GTA 18 12 P30w8 A GCA GCA GAT GAT ACA GTA TT 19 13 P30w9 GCA GCA GAT GAT ACA GTA TT 19 15 P30m11 A GCA GAT AAT ACA GG 19 16 P30m11 A GCA GAT AAT ACA GTA TT 19	D30w1	ACA								GAA	GAA	10	7
P30W3 A GCA GAT GAT ACA GTA TTA 19 9 P30W4 GGA GCA GAT GAT ACA GTA TTA 20 10 P30W5 GGA GCA GAT GAT ACA GTA TTA 21 11 P30W6 ACA GGA GCA GAT GAT ACA 18 12 P30W7 CA GGA GCA GAT GAT ACA GT 19 13 P30W8 A GGA GCA GAT GAT ACA GT 19 15 P30W10 ACA GGA GCA GAT AAT ACA GGT 19 16 P30m11 A GCA GAT AAT ACA GTA TT 19 18 P30m12 GA GCA GAT AAT ACA GTA TT 19 18	•												
P30w4 GGA GCA GAT GAT GAC GTA TTA 20 10 P30w6 ACA GGA GCA GAT GAT ACA GTA TTA 21 11 P30w7 CA GGA GCA GAT GAT ACA GT 19 13 P30w8 A GGA GCA GAT GAT ACA GT 19 15 P30w10 ACA GGA GCA GAT GAT ACA GT 19 16 P30m11 A GCA GAT AAT ACA GTA TT 18 17 P30m12 GA GCA GAT AAT ACA GTA TT 19 18 P30m12 GA GCA GAT AAT ACA GTA TT 19 19 P30m13 A GGA GCA GAT AAT ACA GTA TT													
P30w5 GGA GCA GAT GAT ACA GTA TTA 21 11 P30w6 ACA GGA GCA GAT GAT ACA 18 12 P30w7 CA GGA GCA GAT GAT ACA GT 19 13 P30w8 A GGA GCA GAT GAT ACA GT 19 15 P30w10 ACA GGA GCA GAT GAT ACA GG 19 16 P30m11 A GCA GAT AAT ACA GTA TT 18 17 P30m12 GA GCA GAT AAT ACA GTA TT 19 18 P30m13 A GCA GAT AAT ACA GTA TT 19 19 P30m14 GGA GCA GAT AAT ACA GTA TT 19 23													
P30w6 ACA GGA GCA GAT GAT ACA 18 12 P30w7 CA GGA GCA GAT GAT ACA GT 19 13 P30w8 A GGA GCA GAT GAT ACA GTA TG 20 14 P30w9 GGA GCA GAT GAT ACA GTA TG 19 15 P30w10 ACA GGA GCA GAT GAT ACA GG 19 16 P30m11 A GCA GAT AAT ACA GTA TT 18 17 P30m12 GA GCA GAT AAT ACA GTA TT 19 18 P30m13 A GCA GAT AAT ACA GTA TT 19 18 P30m14 GGA GCA GAT AAT ACA GTA TT 19 19 P30m15 GGA GCA GAT AAT ACA GTA TT 20 20 P30m15 ACA GGA GCA GAT AAT ACA GTA TT 21 21 P30m17 CA GGA GCA GAT AAT ACA GTA TG 19 23 P30m18 A GGA GCA GAT AAT ACA GTA TG 19 23 P30m19 GGA GCA GAT AAT ACA GTA TG 19 25 P30m20 ACA GGA GCA GAT AAT ACA GTA GT 19 25 P30m21 A GCA GAT GAT AAT ACA GTA GT 19 26			CCA	CCA	CAM	CAI	ACA	GTA	· T.T.				
P30w7 CA GGA GCA GAT GAT ACA GT 19 13 P30w8 A GGA GCA GAT GAT ACA GTA TG 20 14 P30w9 GGA GCA GAT GAT ACA GTA TG 19 15 P30w10 ACA GGA GCA GAT GAT ACA GG 19 16 P30m11 A GCA GAT AAT ACA GG 19 16 P30m11 A GCA GAT AAT ACA GTA TT 18 17 P30m12 GA GCA GAT AAT ACA GTA TT 19 18 P30m13 A GCA GAT AAT ACA GTA TT 19 18 P30m14 GGA GCA GAT AAT ACA GTA TT 19 18 P30m15 GGA GCA GAT AAT ACA GTA TT 20 20 P30m15 ACA GGA GCA GAT AAT ACA 18 22 P30m16 ACA GGA GCA GAT AAT ACA 18 22 P30m17 CA GGA GCA GAT AAT ACA GTA TG 19 23 P30m20 ACA GGA GCA GAT AAT ACA GTA TG 19 25 P30m20 ACA GGA GCA GAT AAT ACA GTA TG 19 26 P30m21 A GCA GAT GAT AAT ACA GTA GT 15 27 <		202	GGA	GCA	GAT	GAT	ACA	GTA	TTA				
P30w8 A GGA GCA GAT GAT ACA GTA TG 20 14 P30w9 GGA GCA GAT GAT ACA GTA TG 19 15 P30w10 ACA GGA GCA GAT GAT ACA GG 19 16 P30w11 A GCA GCA GAT AAT ACA GTA TT 18 17 P30m12 GA GCA GAT AAT ACA GTA TT 18 17 P30m12 GA GCA GAT AAT ACA GTA TT 19 18 P30m13 A GCA GCA GAT AAT ACA GTA TT 19 18 P30m14 GGA GCA GAT AAT ACA GTA TT 19 19 P30m15 GGA GCA GAT AAT ACA GTA TT 20 20 P30m15 ACA GGA GCA GAT AAT ACA GTA TTA 18 22 P30m17 CA GGA GCA GAT AAT ACA GTA TTA 19 23 P30m18 A GGA GCA GAT AAT ACA GTA TG 19 23 P30m20 ACA GGA GCA GAT AAT ACA GTA TG 19 25 P30m20 ACA GGA GCA GAT AAT ACA GTA TG 19 26 P30m21 A GCA GAT GAT AAT ACA GTA GT 15 27 P30m22 A GCA GAT GAT AAT ACA GTA GT 15 29 <td></td>													
P30w9 GGA GCA GAT GAT ACA GTA TG 19 15 P30w10 ACA GGA GCA GAT GAT GAT ACA GG 19 16 P30m11 A GCA GAT AAT ACA GTA TT 18 17 P30m12 GA GCA GAT AAT ACA GTA TT 19 18 P30m13 A GCA GAT AAT ACA GTA TT 19 19 P30m14 GGA GCA GAT AAT ACA GTA TT 20 20 P30m15 GCA GCA GCA AAT ACA GTA 18 22 P30m17 CA GGA GCA GAT AAT ACA GT 19 23 P30m18 A GGA GCA GAT AAT ACA GT 19 25													
P30w10 ACA GGA GCA GAT GAT ACA GG 19 16 P30m11 A GCA GAT AAT ACA GTA TT 18 17 P30m12 GA GCA GAT AAT ACA GTA TT 19 18 P30m13 A GCA GAT AAT ACA GTA TT 19 19 P30m14 GGA GCA GAT AAT ACA GTA TT 20 20 P30m15 GGA GCA GAT AAT ACA GTA TT 20 20 P30m15 CA GGA GCA GAT AAT ACA GTA TT 19 23 P30m17 CA GGA GCA GAT AAT ACA GT 19 26 P30m18 A GGA GCA GAT AAT ACA GT TG		Α											
P30m11 A GCA GAT AAT ACA GTA TT 18 17 P30m12 GA GCA GAT AAT ACA GTA TT 19 18 P30m13 A GCA GAT AAT ACA GTA TT 19 19 P30m14 GGA GCA GAT AAT ACA GTA TT 20 20 P30m15 GGA GCA GAT AAT ACA GTA TT 20 20 P30m15 ACA GGA GCA GAT AAT ACA GTA TT 21 21 21 21 21 21 21 21 21 21 21 21 22 20 24 20 24 23 22 24 23 23 22 24 23 23 22 24 24 23 22 24 24 23 22									TG	•			
P30m12 GA GCA GAT AAT ACA GTA TT 19 18 P30m13 A GCA GAT AAT ACA GTA TTA 19 19 P30m14 GGA GCA GAT AAT ACA GTA TT 20 20 P30m15 GGA GCA GAT AAT ACA GTA TT 21 21 P30m15 ACA GGA GCA GAT AAT ACA TTA 21 21 21 21 21 21 21 21 21 21 22 22 22 23 23 23 23 23 23 23 23 23 23 23 23 24 23 23 24 23 24 23 24 23 23 24 24 23 24 24 23 22 24 24 24 24 24 <t< td=""><td></td><td>ACA</td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></t<>		ACA											
P30m13 A GCA GAT AAT ACA GTA TTA 19 19 P30m14 GGA GCA GAT AAT ACA GTA TTT 20 20 P30m15 GGA GCA GAT AAT ACA GTA TTA 21 21 P30m15 ACA GGA GCA GAT AAT ACA GTA TTA 20 24 P30m17 CA GGA GCA GAT AAT ACA GT 19 23 P30m18 A GGA GCA GAT AAT ACA GT 19 25 P30m20 ACA GGA GCA GAT AAT ACA GT 19 26 P30w21 A GCA GAT AAT ACA GT 19 25 P30m21 A GCA GAT GAT ACA GT 15 27 29													
P30m14 GGA GCA GAT AAT ACA GTA TTT 20 20 P30m15 GGA GCA GAT AAT ACA GTA TTA 21 21 P30m15 ACA GGA GCA GAT AAT ACA TTA 21 21 P30m15 ACA GGA GCA GAT AAT ACA TTA 18 22 P30m18 A GGA GCA GAT AAT ACA GT 19 23 P30m18 A GGA GCA GAT AAT ACA GTA 19 25 P30m19 GGA GCA GAT AAT ACA GTA 19 25 P30m20 ACA GGA GAT AAT ACA GT 19 26 P30w21 A GCA GAT AAT ACA GTA GT 16 28 P30w22 A													18
P30m15 GGA GCA GAT AAT ACA GTA TTA 21 21 P30m15 ACA GGA GCA GAT AAT ACA 18 22 P30m17 CA GGA GCA GAT AAT ACA GT 19 23 P30m19 GGA GCA GAT AAT ACA GTA TG 19 25 P30m20 ACA GGA GCA GAT AAT ACA GG 19 26 P30m20 ACA GGA GAT AAT ACA GG 19 26 P30m21 A GCA GAT GAT ACA GT 15 27 27 23 26 28 28 20 24 26 28 26 28 29 26 28 26 26 26 16 28 29 20 26 26 27 27 20<													19
P30m15 ACA GGA GCA GAT AAT ACA 18 22 P30m17 CA GGA GCA GAT AAT ACA GT 19 23 P30m18 A GGA GCA GAT AAT ACA GTA TG 19 25 P30m19 GGA GCA GAT AAT ACA GTA TG 19 26 P30m20 ACA GGA GCA GAT AAT ACA GT 19 26 P30w21 A GCA GAT AAT ACA GT 15 27 19 26 20 24 20 24 25 22 24 25 26 26 26 16 28 25 26 26 26 16 28 26 26 26 26 26 26 26 26 26 26 26 26 26 26												20	20
P30m17 CA GGA GCA GAT AAT ACA GT 19 23 P30m18 A GGA GCA GAT AAT ACA GTA TG 20 24 P30m19 GGA GCA GAT AAT ACA GTA TG 19 25 P30m20 ACA GGA GCA GAT AAT ACA GT 19 26 p30w21 A GCA GAT GAT ACA GT 15 27 p30w22 A GCA GAT GAT AAT ACA GTA 15 29 p30m23 A GCA GAT AAT ACA GTA 15 29 p30m24 A GCA GAT GAT AAT ACA GTA 15 29 p30w25 GCA GAT GAT ACA GT 14 31 p30w26 A GCA GAT GAT ACA GT 14 32 p30w27 CA GAT GAT ACA GT 13 33 p30w28 GA GCG GAT GAT ACA 13 35 p30m30 GCA GAT GAT ACA 13 35 p30m31 GCA GAT GAT ACA GT 14 37 p30w32 GCA GAT GAC ACA GT 14 38 p30w34 CA GAT GAC ACA GT 14 39 p30w35 GCA GAT GAT ACA ATA TG								GTA	TTA	•		21	21
P30m18 A GGA GCA GAT AAT ACA GTA TG 20 24 P30m19 GGA GCA GAT AAT ACA GTA TG 19 25 P30m20 ACA GGA GCA GAT AAT ACA GG 19 26 p30w21 A GCA GAT GAT ACA GT 15 27 p30w22 A GCA GAT GAT ACA GTA G 16 28 p30m23 A GCA GAT AAT ACA GTA G 15 29 p30m24 A GCA GAT AAT ACA GTA G 16 30 p30w25 GCA GAT GAT ACA GT 14 31 p30w26 A GCA GAT GAT ACA GT 14 32 p30w27 CA GAT GAT ACA GT 13 33 p30w28 GA GCG GAT GAT ACA 14 34 p30w29 A GCG GAT GAT ACA 13 35 p30m30 GCA GAT AAT ACA GTA 15 36 p30m31 GCA GAT GAC ACA GT 14 37 p30w32 GCA GAT GAC ACA GT 14 38 p30w34 CA GAT GAC ACA GT 14 39 p30w35 GCA GAC GAT ACA ATA TG												18	22
P30m19 GGA GCA GAT AAT ACA GTA TG 19 25 P30m20 ACA GGA GCA GAT AAT ACA GG 19 26 p30w21 A GCA GAT ACA GT 15 27 p30w22 A GCA GAT GAT ACA GTA GT 16 28 p30w22 A GCA GAT GAT ACA GTA GT 16 28 p30m23 A GCA GAT AAT ACA GTA 15 29 p30m24 A GCA GAT AAT ACA GTA GT 16 30 p30w25 GCA GAT GAT ACA GT 14 31 31 32 33 33 33 33 33 33 33 33 33 33 33 34 34 34 34 </td <td>P30m17</td> <td></td> <td>19</td> <td>23</td>	P30m17											19	23
P30m20 ACA GGA GCA GAT AAT ACA GG 19 26 p30w21 A GCA GAT GAT ACA GT 15 27 p30w22 A GCA GAT GAT ACA GTA G 16 28 p30w23 A GCA GAT AAT ACA GTA 15 29 p30m24 A GCA GAT AAT ACA GTA 16 30 p30w25 GCA GAT GAT ACA GT 14 31 p30w26 A GCA GAT GAT ACA GG 14 32 p30w27 CA GAT GAT ACA GT 13 33 p30w28 GA GCG GAT GAT ACA 14 34 p30w29 A GCG GAT ACA GTA 15 36 p30m31 GCA GAT GAT ACA GT 14 37		Α	GGA	GCA	GAT	AAT	ACA	GTA	TG			20	24
p30w21 A GCA GAT GAT ACA GT 15 27 p30w22 A GCA GAT GAT ACA GTA G 16 28 p30m23 A GCA GAT AAT ACA GTA 15 29 p30m24 A GCA GAT AAT ACA GTA G 16 30 p30w25 GCA GAT GAT ACA GT 14 31 p30w26 A GCA GAT GAT ACA GG 14 32 p30w27 CA GAT GAT ACA GT 13 33 p30w28 GA GCG GAT GAT ACA 14 34 p30w29 A GCG GAT GAT ACA 13 35 p30m30 GCA GAT AAT ACA GTA 15 36 p30m31 GCA GAT AAT ACA GT 14 37 p30w32 GCA GAT GAC ACA GT 14 38 p30w33 CA GAT GAC ACA GTA G 14 39 p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA GT 14 43 p30w36 GCA GAC GAT ACA GT 14 43 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44	P30m19		GGA	GCA	GAT	AAT	ACA	GTA	TG			19	25
p30w22 A GCA GAT GAT ACA GTA G 16 28 p30m23 A GCA GAT AAT ACA GTA 15 29 p30m24 A GCA GAT AAT ACA GTA G 16 30 p30w25 GCA GAT GAT ACA GT 14 31 p30w26 A GCA GAT GAT ACA GG 14 32 p30w27 CA GAT GAT ACA GT 13 33 p30w28 GA GCG GAT GAT ACA 14 34 p30w29 A GCG GAT GAT ACA 13 35 p30m30 GCA GAT AAT ACA GTA 15 36 p30m31 GCA GAT AAT ACA GT 14 37 p30w32 GCA GAT GAC ACA GT 14 38 p30w33 CA GAT GAC ACA GT 14 39 p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GT 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45	P30m20	ACA	GGA	GCA	GAT	AAT	ACA	GG.		•		19	26
p30w22 A GCA GAT GAT ACA GTA G 16 28 p30m23 A GCA GAT AAT ACA GTA 15 29 p30m24 A GCA GAT AAT ACA GTA G 16 30 p30w25 GCA GAT GAT ACA GT 14 31 p30w26 A GCA GAT GAT ACA GG 14 32 p30w27 CA GAT GAT ACA GT 13 33 p30w28 GA GCG GAT GAT ACA 14 34 p30w29 A GCG GAT GAT ACA 13 35 p30m30 GCA GAT AAT ACA GTA 15 36 p30m31 GCA GAT AAT ACA GT 14 37 p30w32 GCA GAT GAC ACA GT 14 38 p30w33 CA GAT GAC ACA GT 14 39 p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GT 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45	p30w21		A	GCA	GAT	GAT	ACA	GT				15	27
p30m23 A GCA GAT AAT ACA GTA 15 29 p30m24 A GCA GAT AAT ACA GTA G 16 30 p30w25 GCA GAT GAT ACA GT 14 31 p30w26 A GCA GAT GAT ACA GG 14 32 p30w27 CA GAT GAT ACA GT 13 33 p30w28 GA GCG GAT GAT ACA 14 34 p30w29 A GCG GAT GAT ACA 13 35 p30m30 GCA GAT AAT ACA GTA 15 36 p30m31 GCA GAT AAT ACA GT 14 37 p30w32 GCA GAT GAC ACA GT 14 38 p30w33 CA GAT GAC ACA GT 14 39 p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GT 14 43 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45	p30w22		Α	GCA	GAT	GAT	ACA	GTA	G			16	
p30w25 GCA GAT GAT ACA GT 14 31 p30w26 A GCA GAT GAT ACA GG 14 32 p30w27 CA GAT GAT ACA GT 13 33 p30w28 GA GCG GAT GAT ACA 14 34 p30w29 A GCG GAT GAT ACA 13 35 p30m30 GCA GAT AAT ACA GTA 15 36 p30m31 GCA GAT AAT ACA GT 14 37 p30w32 GCA GAT GAC ACA GT 14 38 p30w33 CA GAT GAC ACA GTA G 14 39 p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GT 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45	p30m23		Α	GCA	GAT	AAT	ACA	GTA				15	
p30w25 GCA GAT GAT ACA GT 14 31 p30w26 A GCA GAT GAT ACA GG 14 32 p30w27 CA GAT GAT ACA GT 13 33 p30w28 GA GCG GAT GAT ACA 14 34 p30w29 A GCG GAT GAT ACA 13 35 p30m30 GCA GAT AAT ACA GTA 15 36 p30m31 GCA GAT AAT ACA GT 14 37 p30w32 GCA GAT GAC ACA GT 14 38 p30w33 CA GAT GAC ACA GTA G 14 39 p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GT 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45	p30m24		Α	GCA	GAT	AAT	ACA	GTA	G				
p30w26 A GCA GAT GAT ACA GG 14 32 p30w27 CA GAT GAT ACA GT 13 33 p30w28 GA GCG GAT GAT ACA 14 34 p30w29 A GCG GAT GAT ACA 13 35 p30m30 GCA GAT AAT ACA GTA 15 36 p30m31 GCA GAT AAT ACA GT 14 37 p30w32 GCA GAT GAC ACA GT 14 38 p30w33 CA GAT GAC ACA GTA G 14 39 p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GT 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45				GCA	GAT	GAT	ACA	GT				14	
p30w27 CA GAT GAT ACA GT 13 33 p30w28 GA GCG GAT GAT ACA 14 34 p30w29 A GCG GAT GAT ACA 13 35 p30m30 GCA GAT AAT ACA GTA 15 36 p30m31 GCA GAT AAT ACA GT 14 37 p30w32 GCA GAT GAC ACA GT 14 38 p30w33 CA GAT GAC ACA GTA G 14 39 p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GG 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45			Α	GCA	GAT	GAT	ACA	GG				14	
p30w28 GA GCG GAT GAT ACA 14 34 p30w29 A GCG GAT GAT ACA 13 35 p30m30 GCA GAT AAT ACA GTA 15 36 p30m31 GCA GAT AAT ACA GT 14 37 p30w32 GCA GAT GAC ACA GT 14 38 p30w33 CA GAT GAC ACA GTA G 14 39 p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GG 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45			•	CA	GAT	GAT	ACA	GT				13	
p30w29 A GCG GAT GAT ACA 13 35 p30m30 GCA GAT AAT ACA GTA 15 36 p30m31 GCA GAT AAT ACA GT 14 37 p30w32 GCA GAT GAC ACA GT 14 38 p30w33 CA GAT GAC ACA GTA G 14 39 p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GG 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45			GA	GCG	GAT	GAT	ACA					14	
p30m30 GCA GAT AAT ACA GTA 15 36 p30m31 GCA GAT AAT ACA GT 14 37 p30w32 GCA GAT GAC ACA GT 14 38 p30w33 CA GAT GAC ACA GTA G 14 39 p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GG 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45			Α	GCG	GAT	GAT	ACA					13	
p30m31 GCA GAT AAT ACA GT 14 37 p30w32 GCA GAT GAC ACA GT 14 38 p30w33 CA GAT GAC ACA GTA G 14 39 p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GG 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45				GCA	GAT	AAT	ACA	GTA					
p30w32 GCA GAT GAC ACA GT 14 38 p30w33 CA GAT GAC ACA GTA G 14 39 p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GG 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45	p30m31			GCA	GAT	AAT	ACA	GT					
p30w33 CA GAT GAC ACA GTA G 14 39 p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GG 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45													
p30w34 CA GAT GAT ACA ATA TT 16 40 p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GG 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45				CA	GAT	GAC	ACA	GTA	G				
p30w35 GCA GAT GAT ACA ATA TG 16 41 p30w36 GCA GAC GAT ACA GG 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45													
p30w36 GCA GAC GAT ACA GG 13 42 p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45													
p30w37 GCA GAC GAT ACA GT 14 43 p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45													
p30w38 A GAT GAT ACA ATA TT 15 44 p30w39 A GAT GAT ACA ATA TTA 16 45													
p30w39 A GAT GAT ACA ATA TTA 16 45									TT				
p30w40 GCA GAT GAT ACA ATA 15 46	p30w40											15	46

Table 3 - Cont'd-1

	44	45	46	47	48	49	50	51	52	53	54	length	Sea	מד
		AAA			GGG				GGT				Deq	
P48w1	0011		1110		GGG			GGA		GG	1110	. 18	47	
P48w2					GGG			GGA		TG		19	48	
P48w3					GGG			GGA		TTG		20	49	
					GGG			GGA					50	
P48w4			_							TTT		21		
P48w5									GGT	TTG		21	. 51	
P48w6					GGG				_			18	52	
P48,w7		_							G			19	53	
P48w8					GGG							19	54	
P48w9					GGG							20	55	
P48w10		, A			GGG				GGG	GG		22	56	
P48w21					GGG			GGA				18	57	•
P48w22					GGG							18	58	
P48w23					GGG			GGA				19	59	
P48w24		Α	ATG		GGG			GGA				19	60	
P48w25				ATA	GGG	GGA	ATT	GGA	GGT	GG		18	61	
P48w26				ATA	GGG	GGA	TTA	GGA	GGT	TG		19	62	
P48w28				ATA	GGG	GGA	ATT	GGA	GGT	TTG		20	63	
P48w29				ATA	GGG	GGA	ATT	GGA	GGT	TTT		21	64	
P48m11				GTA	GTG	GGA	ATT	GGA	GGT	GG		18	65	
P48m12				GTA	GTG	GGA	ATT	GGA	GGT	TG		19	66	
P48m13					GTG			GGA		TTG		20	67	
P48m14					GTG					TTT		21	68	
P48m15			G						GGT			21	69	
P48m16					GTG			GGA				18	70	
P48m17	•				GTG				G			19	71	
P48m18		· A			GTG			GGA	_			19	72	
P48m19					GTG				G			20	73	
P48m20									GGG	GG		22	74	
P48m29		••	0						GGT			18	75	
P48m30					GTG				GGT			19	76	
P48m31			ΔTG		GTG			GGA	001	+0	•	18	77	
P48m32					GTG			GGA	G			19	78	
P48m33		Δ			GTG			GGA	•			19	79	
p48w34		А			GGG			G				14	80	
p48w35					GGG			Ğ				15	81	
p48w36					GGG			GG				16	82	
p48w37				_	GGG			GG				15	83	
p48m38					GTG			G				14	84	
				•	GTG			G				15	85	
p48m39					GTG			GG				16		
p48m40			TG					GG					86	
p48m41					GTG GGG							15 15	87	
p48w42												15	88	
p48w43					GGG			_				14	89	
p48w44		75			GGG			G				14	90	
p48w45		A			GGA							16	. 91	
p48w46		71 75 75			GGG		ATT					15	92	
p48w47	-				GGG							15	93	
p48w48	A	AAA	ATG	ATA	GGG	GG						15	94	

Table 3 - Cont'd-2

p48w49		AA	ATG	ATA	GGG	GGA	AG			15	95
p48w50		AAA	ATA	ATA	GGG	GGA	AG			16	96
p48w51		AAA	ATA	AAA	AΤ					15	97
p48m52		AAA	ATG	ATA	GTG	GGA	AG			16	98
p48w52b		AAA	TTG	ATA	GGG	GG				14	99
p48m53		AAA	ATG	ATA	GTG	GGA				15	100
p48w53b		AAA	TTG	ATA	GGG	GGA				15	101
p48w54	CA	AAA	TTG	ATA	G					15	102
p48w55			ATG	GTA	GGG	GGA	ATT			15	103
p48w56		AA	ATG	GTA	GGG	GGA				14	104
p48w57	Α	AAA	ATG	GTA	GGG	G				14	105
p48w58			ATG	ATA	GGG	GAA	ATT			15	106
p48w59				ATA	GGG	GAA	ATT	GGA		15	107
p48w60						GAA		GGA	G	16	108
p48w61			ATG	ATA	GGG	GGG	ATT			15	109
p48w62				ATA	GGG			GG		14	110
p48w63				Α	GGG	GGG	ATT	GGA		13	111
p48m64				ATA		GGA				15	112
p48m65				ATA		GGA				16	113
p48m66	CA	AAA		ATA		GG				16	114
p48m67		AAA		ATA		GGA				15	115
p48m68		AAA		ATA		GGA				16	116
p48m69	CA	AAA		ATA		G				15	117
p48w70				ATA		GG				14	118
p48w71				ATA		G				14	119
pc48w72	A	AAA	ATA	ATA	GGG	GGA				16	120

Table 3 - Cont'd-3

P50w2 A GGG GGA ATT GGA GGT TTT 19 P50w3 TA GGG GGA ATT GGA GGT TTT 20 P50w4 A GGG GGA ATT GGA GGT TTT AG 20 P50w5 TA GGG GGA ATT GGA GGT TTT AG 21 P50w6 GTA GGG GGA ATT GGA GGT TGG 19 P50w7 G GTA GGG GGA ATT GGA GGT TGG 20 P50w8 GTA GGG GGA ATT GGA GGT TTG 20 P50w9 GTA GGG GGA ATT GGA GGT TTT 20 P50w10 TG GTA GGG GGA ATT GGA GGT TTT 20 P50w21 GG GGA ATT GGA GGT TTT 17 P50w22 GG GGA ATT GGA GGT TTG 16 P50w23 GG GGA ATT GGA GGT TTT 16 P50w24 GG GGA ATT GGA GGT TTT 15 P50w25 G GGA ATT GGA GGT TTT 17 P50w26 GG GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	q ID 121 122 123 124 125 126 127
P50w1 GGG GGA ATT GGA GGT TTT 18 P50w2 A GGG GGA ATT GGA GGT TTT 19 P50w3 TA GGG GGA ATT GGA GGT TTT 20 P50w4 A GGG GGA ATT GGA GGT TTT AG 20 P50w5 TA GGG GGA ATT GGA GGT TTT AG 21 P50w6 GTA GGG GGA ATT GGA GGT TGG 19 P50w7 G GTA GGG GGA ATT GGA GGT TGG 20 P50w8 GTA GGG GGA ATT GGA GGT TTT 20 P50w9 GTA GGG GGA ATT GGA GGT TTT 17 P50w21 GG GGA ATT GGA GGT	122 123 124 125 126
P50w2 A GGG GGA ATT GGA GGT TTT 19 P50w3 TA GGG GGA ATT GGA GGT TTT 20 P50w4 A GGG GGA ATT GGA GGT TTT AG 20 P50w5 TA GGG GGA ATT GGA GGT TTT AG 21 P50w6 GTA GGG GGA ATT GGA GGT TGG 19 P50w7 G GTA GGG GGA ATT GGA GGT TGG 20 P50w8 GTA GGG GGA ATT GGA GGT TTG 20 P50w9 GTA GGG GGA ATT GGA GGT TTT 20 P50w10 TG GTA GGG GGA ATT GGA GGT TTT 20 P50w21 GG GGA ATT GGA GGT TTT 17 P50w22 GG GGA ATT GGA GGT TTG 16 P50w23 GG GGA ATT GGA GGT TTT 16 P50w24 GG GGA ATT GGA GGT TTT 15 P50w25 G GGA ATT GGA GGT TTT 17 P50w26 GG GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	122 123 124 125 126
P50w3 TA GGG GGA ATT GGA GGT TTT 20 P50w4 A GGG GGA ATT GGA GGT TTT AG 20 P50w5 TA GGG GGA ATT GGA GGT TTT AG 21 P50w6 GTA GGG GGA ATT GGA GGT TGG 19 P50w7 G GTA GGG GGA ATT GGA GGT TGG 20 P50w8 GTA GGG GGA ATT GGA GGT TTG 20 P50w9 GTA GGG GGA ATT GGA GGT TTT 20 P50w10 TG GTA GGG GGA ATT GGA GGT TTT 20 P50w21 GG GGA ATT GGA GGT TTT 17 P50w22 GG GGA ATT GGA GGT TTG 16 P50w23 GG GGA ATT GGA GGT TTT AG 18 P50w24 GG GGA ATT GGA GGT TTT AT 18 P50w25 G GGA ATT GGA GGT TTT AT 18 P50w26 G GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	123 124 125 126
P50w4 A GGG GGA ATT GGA GGT TTT AG 20 P50w5 TA GGG GGA ATT GGA GGT TTT AG 21 P50w6 GTA GGG GGA ATT GGA GGT TGG 19 P50w7 G GTA GGG GGA ATT GGA GGT TGG 20 P50w8 GTA GGG GGA ATT GGA GGT TTG 20 P50w9 GTA GGG GGA ATT GGA GGT TTT 20 P50w10 TG GTA GGG GGA ATT GGA GGT GG 20 P50w21 GG GGA ATT GGA GGT TTT 17 P50w22 GG GGA ATT GGA GGT TTG 16 P50w23 GG GGA ATT GGA GGT TTT AG 18 P50w24 GG GGA ATT GGA GGT TTT AT 18 P50w25 G GGA ATT GGA GGT TTT AT 18 P50w26 GG GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	124 125 126
P50w5 TA GGG GGA ATT GGA GGT TTT AG 21 P50w6 GTA GGG GGA ATT GGA GGT TGG 19 P50w7 G GTA GGG GGA ATT GGA GGT TGG 20 P50w8 GTA GGG GGA ATT GGA GGT TTG 20 P50w9 GTA GGG GGA ATT GGA GGT TTT 20 P50w10 TG GTA GGG GGA ATT GGA GGT GG 20 P50w21 GG GGA ATT GGA GGT TTT 17 P50w22 GG GGA ATT GGA GGT TTG 16 P50w23 GG GGA ATT GGA GGT TTT 16 P50w24 GG GGA ATT GGA GGT TTT 15 P50w25 G GGA ATT GGA GGT TTT 17 P50w26 GG GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	125 126
P50w6 GTA GGG GGA ATT GGA GGT TGG 19 P50w7 G GTA GGG GGA ATT GGA GGT TGG 20 P50w8 GTA GGG GGA ATT GGA GGT TTG 20 P50w9 GTA GGG GGA ATT GGA GGT TTT 20 P50w10 TG GTA GGG GGA ATT GGA GGT GG 20 P50w21 GG GGA ATT GGA GGT TTT 17 P50w22 GG GGA ATT GGA GGT TTG 16 P50w23 GG GGA ATT GGA GGT TTT AG 18 P50w24 GG GGA ATT GGA GGT TT 15 P50w25 G GGA ATT GGA GGT TT 17 P50w26 GG GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	126
P50w7 G GTA GGG GGA ATT GGA GGT TGG 20 P50w8 GTA GGG GGA ATT GGA GGT TTG 20 P50w9 GTA GGG GGA ATT GGA GGT TTT 20 P50w10 TG GTA GGG GGA ATT GGA GGT GG 20 p50w21 GG GGA ATT GGA GGT TTT 17 P50w22 GG GGA ATT GGA GGT TTG 16 P50w23 GG GGA ATT GGA GGT TTT AG 18 P50w24 GG GGA ATT GGA GGT TT 15 P50w25 G GGA ATT GGA GGT TT 17 P50w26 GG GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	
P50w8 GTA GGG GGA ATT GGA GGT TTG 20 P50w9 GTA GGG GGA ATT GGA GGT TTT 20 P50w10 TG GTA GGG GGA ATT GGA GGT GG 20 P50w21 GG GGA ATT GGA GGT TTT 17 P50w22 GG GGA ATT GGA GGT TTG 16 P50w23 GG GGA ATT GGA GGT TTT AG 18 P50w24 GG GGA ATT GGA GGT TT 15 P50w25 G GGA ATT GGA GGT TTT 17 P50w26 GG GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	127
P50w9 GTA GGG GGA ATT GGA GGT TTT 20 P50w10 TG GTA GGG GGA ATT GGA GGT GG 20 p50w21 GG GGA ATT GGA GGT TTT 17 P50w22 GG GGA ATT GGA GGT TTG 16 P50w23 GG GGA ATT GGA GGT TTT AG 18 P50w24 GG GGA ATT GGA GGT TT 15 P50w25 G GGA ATT GGA GGT TTT AT 18 P50w26 GG GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	
P50w10 TG GTA GGG GGA ATT GGA GGT GG 20 p50w21 GG GGA ATT GGA GGT TTT 17 P50w22 GG GGA ATT GGA GGT TTG 16 P50w23 GG GGA ATT GGA GGT TTT AG 18 P50w24 GG GGA ATT GGA GGT TG 15 P50w25 G GGA ATT GGA GGT TTT AT 18 P50w26 GG GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	128
p50w21 GG GGA ATT GGA GGT TTT 17 p50w22 GG GGA ATT GGA GGT TTG 16 p50w23 GG GGA ATT GGA GGT TTT AG 18 p50w24 GG GGA ATT GGA GGT TG 15 p50w25 G GGA ATT GGA GGT TTT AT 18 p50w26 GG GGA ATT GGA GGT TTT 17 p50m11 GGG GGA GTT GGA GGT TTT 18 p50m12 A GGG GGA GTT GGA GGT TTT 19 p50m13 TA GGG GGA GTT GGA GGT TTT 20	129
P50w22 GG GGA ATT GGA GGT TTG 16 P50w23 GG GGA ATT GGA GGT TTT AG 18 P50w24 GG GGA ATT GGA GGT TTG 15 P50w25 G GGA ATT GGA GGT TTT AT 18 P50w26 GG GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	130
P50w23 GG GGA ATT GGA GGT TTT AG 18 P50w24 GG GGA ATT GGA GGT TG 15 P50w25 G GGA ATT GGA GGT TTT AT 18 P50w26 GG GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	131
P50w24 GG GGA ATT GGA GGT TG 15 P50w25 G GGA ATT GGA GGT TTT AT 18 P50w26 GG GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	132
P50w25 G GGA ATT GGA GGT TTT AT 18 P50w26 GG GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	133
P50w26 GG GGA ATT GGA GGT TTT 17 P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	134
P50m11 GGG GGA GTT GGA GGT TTT 18 P50m12 A GGG GGA GTT GGA GGT TTT 19 P50m13 TA GGG GGA GTT GGA GGT TTT 20	135
P50m12 A GGG GGA GTT GGA GGT TTT 1.9 P50m13 TA GGG GGA GTT GGA GGT TTT 20	136
P50m13 . TA GGG GGA GTT GGA GGT TTT 20	137
200120	138
	139
P50m14 A GGG GGA GTT GGA GGT TTT AG 20	140
100111120	141
100m20	142
P50m17 G GTA GGG GGA GTT GGA GGT TGG 20	143
2001120	144
10011129	145
	146
LOVINE	147
2 0 0 11.2 0	148
	149
	150
poone	151
Poor of	152
Formor	153
F-0-1110-1	154
p	155
p come c	156
g	
F	157
p50w39 GA ATT GGG GGT TTT 14	

p50w40		GA	ATT	GGG	GGT	TTT	AG	15	160
p50w41		GGA	ATT	GGG	GGT	TG		13	161
p50w42		GGA	ATT	GGG	GGT	G		12	162
p50w43		GA	ATT	GGG	GGT	TG		12	163
p50w44		GA	ATT	GGG	GGT	TTG		13	164
p50w45	GGG	GGA	ATT	GCA	G			13	165
p50w46		GGA	ATT	GCA	GGT	TG		14	166
p50w47		GGA	ATT	GCA	GGT	Ģ		13	167
p50w48		GGA	ATT	GGA	GGG	TTG		14	168
p50w49		GA	ATT	GGA	GGG	TTG		13	169
p50w50		GA	ATT	GGA	GGG	TTT		14	170
p50w51		GGA	ATT	GGA	GGC	TTG		14	171
p50w52		GA	ATT	GGA	GGC	TTG		13	172
p50w53		GA	ATT	GGA	GGC	TTT		14	173
p50m54		GGA	GTT	GGA	GGT	TTG		15	174
p50m55		GA	GTT	GGA	GGT	TTT		14	175

Table 3 - Cont'd-5

•	51	52	53	54	55	56	57	58	length	Seq ID
	GGA	GGT	TTT	ATC	AAA	GTA	AGA	CAG		
p54w1		GGT	TTT	ATC	AAA	GTA	A		16	176
p54w2		GT	TTT	ATC	AAA	GTA	AG		16	177
p54w3		GT	TTT	ATC	AAA	GTA	AGA		17	178
p54w4		T	TTT	ATC	AAA	GTA	AGA		16	179
p54w5		GGT	TTT		AAA				15	180
p54w6		GT	TTT		AAA				15	181
p54m7		GGT	TTT		AAA			3	15	182
p54m7		GT	TTT		AAA		Ά	-	15	183
p54m0 p54m9	•	GT	TTT		AAA				16	184
p54m3		T	TTT		AAA				16	185
		GGT	TTT		AAA		non		14	186
p54m11			TTT		AAA				14	187
p54m12		GT					מל מל		16	188
p54w13		GT	TTT		AAG					
p54w14			TTT		AAG		Α		16	189
p54w15	A	GGT	TTT		AAG				16	190
p54w16		GT	TTT		AAA				17	191
p54w17			TTT		AAA			C	16	192
p54w18		GGC			AAA		Α		17	193
p54w19		GGC			AAA				16	194
p54m20	Α	GGT	TTT	ATT	AAA	GTA	A		17 '	195
p54m21		GGT	TTT	ATT			AG		17	196
p54w22	GA	GGT	TTT	ATT	AAA	GTA			17	197
p54m22	GA	GGT	TTT	ATT	AAA	GTA			17	198
p54m23		GGT	TTT	ATT	GGT	TTT	AT		16	199
p54m24		GGT	TTC	ATT	AAG	GTA			15	200
p54m25		GGT	TTC	ATT	AAG	GTA	Α		16	201
p54w26	Α	GGT	TTC	ATT	AAG	GTA			16	202
p54m26		GGT	TTC	ATT	AAG	GTA			16	203
p54w27		GGT	TTT	ATT		GTA	Α		16	204
p54m27		GGT	TTT	ATT		GTA			16	205
p54m28	Ά	GGT	TTT	ATT		GTA			16	206
p54m29		GGT	TTT	ATT					16	207
p54m30	Gr	GGT	TTT	ATT		GTA	AG		17	208
p54m30 p54w31		GGT	TTT		AAA				16	209
p54w31 p54w32	7\	GGT	TTT		AAA				17	210
p54w32		GGT			AAA		А		16	211
p54w33									16	212
p54w34	GA		TTT		AAA				15	213
p54m35			TTT		AAA				16	213
p54m36			TTT		AAA					
p54m37		GGT	TTT		AGA				15	215
p54m38		GGT			AGA				16	216
p54w39		GGG			AAA				15	217
p54w40		GGG			AAA		Α		16	218
p54w41		GGC			AAA				14	219
p54w42	GA				AAA				14	220
p54m48		GGT			AAA				14	221
p54m49		GT	TTT	GTC	AGA	GTA			14	222

p54m50	GGT	TTT	GTC	AGA	GT	14	223
p54w51	A GGT	TTA	ATC	AAA	GTA	16	224
p54w52	GA GGT					16	225
p54m53				AAA		15	226
p54m54	GGT	TTT	ACC	AAA	GT	14	227

Table 3 - Cont'd-7

		70	~ ~	0.1	~~	00	•					
	78		80		82	83	84	85	86	87		Seq ID
		CCT										
P82w1		CCT	ACA	CCT	GTC	AAC	ATA	AG			19	228
P82w2		CCT	ACA	CCT	GTC	AAC	ATA	ATG			20	229
P82w3	•	CCT	ACA	CCT	GTC	AAC	ATA	ATT			21	230
P82w4	А	CCT	ACA	CCT	GTC	AAC	АТА	AG			20	231
P82w5	7	CCT	DCD.	CCT	CTC	AAC	מידמ	አጥር			21	232
P82w6	Δ 7	CCT	ACA	CCT	CTC	אאכ	עשע	AIG			10	
	A	CCI	ACA	CCI	GIC	AAC	VIV				19	233
P82w7	GA	CCT	ACA	CCT	GTC	AAC	ATA				19 20 21 20 21 19 20	234
P82w8			CA	CCT	GTC	AAC	ATA	ATT	GGA		20	235
P82w9	•		A	CCT	GTC	AAC	ATA	ATT	GGA	A	20	236
P82w10 P82W21 P82m11			ACA	CCT	GTC	AAC	ATA	ATT	GG		20 19	237
P82W21			Α	CCT	GTC	AAC	ATA	ATT	GGA		19	238
P82m11		CCT	ACA	CCT	ACC	AAC	ATA	AG			19	239
P82m12		CCT	ACA	CCT	ACC	AAC	ATA	ATG			20	240
P82m13		CCT	ACA	CCT	ACC	AAC	ATA	ATT	•		21	241
P82m14	Α	CCT	ACA	CCT	ACC	AAC	ATA	AG			19 19 20 21 20 21 19 20	242
P82m15	Δ	CCT	ACA	ССТ	ACC	AAC	ልጥል	ATG			21	243
P82m16	7.	CCT	DCD.	CCT	ACC	אאר	מידמ	1110			10	244
P82m17	$C_{\mathbf{J}}$	CCT	ACA	CCT	ACC.	770	V ILIV				7.9	
	GA	CCI	ACA	CCT	ACC	AAC	VUV	7 mm	CCA		20	245
P82m18												246
P82m19			A	CCT	ACC	AAC	ATA	ATT	GGA	Α	20	247
P82m20			ACA	CCT	ACC	AAC	ATA	ATT	G		19	248
P82m22		CCT	ACA	CCT	TTC	AAC	ATA	ATT			20 19 21 21 21	249
P82m23		CCT	ACA	CCT	GCC	AAC	ATA	ATT			21	250
P82m24		CCT	ACA	CCT	TCC	AAC	ATA	ATT			21	251
P82m25			Α	CCT	TTC	AAC	ATA	ATT	GGA	Α	20	252
P82m26		•	A	CCT	GCC	AAC	ATA	ATT	GGA	A	20	253
P82m27			Α	CCT	TTC	AAC	ATA	ATT	GGA	A	20	254
P82m28			Α	CCT	ACC	AAC	ATA	ATT			16	255
P82m29						AAC			GGA		19	256
P82m30			-	COM	000	770	7.07	70 mm	001		10	257
P82m31			Δ	CCT	TCC	AAC	ΔΤΆ	Δηη	GGA		19	258
P82w32		T	מיא	CCT	GTC	AAC	አ ጥ	7111	COA		15	259
P82w33			ACA	CCT	GTC	AAC	አጥአ ጉ፲				16	
P82w34		1	ACA	CCT	CTC	AAC	V LL V				15	260
	•		ACA	CCT	CTC	AAC AAC AAC AAC AAC AAC	WIW				15	261
P82w35			CA	CCT	GIC	AAC	ATA				14	262
			ACA	CCT	ACC	AAC	ATA				15	263
P82m37			Cii	CCI	1100	11110	*****				7.3	264
P82m38						AAC					15	265
P82m39						AAC					14	266
P82m40						AAC					15	267
P82m41						AAC					14	268
P82w42			CA	CCT	GTC	AAC	GTA				14	269
P82w43						AAC					13	270
P82w44		CCT	ACA								15	271
P82w45	•		ACG				ΑТ				15	272
P82w46			ACG								15	273
P82m47		01				AAC		• •			15	
F0ZIII4/			NCM	CCI	100	MAC	VIV				12	274

P82m48 P82m49 P82m50 P82m51 P82m52 P82m53 P82w54 P82w55 P82w56 P82w57 P82w58 P82w59 P82w60		ACA ACA CA CA A A	CCT CCT CCT CCT CCT CCT CCG CCT CCT	TCC ATC ATC ATC GTC GTT GTC GTC GTC	AAT AAT	ATA ATA ATA ATA ATA ATA ATA ATA ATA	ATG ATT ATT ATG ATT ATT	G	14 14 15 16 15 16 15 16 15 16	275 276 277 278 279 280 281 282 283 284 285 286
P82w61 P82m62 P82m63 P82m64 P82m65 P82m66 P82m67 P82m68 P82m69		CA CA ACA CA	CCT CCT CCT CCT CCT CCT	GCC GCC ACC ACC ACC ACC TTC	AAT AAC AAC AAC AAC AAC AAC	GTA GTA GTA GTA	ATT AG ATT ATG		16 15 15 14 14 14 15	288 289 290 291 292 293 294 295 296
P82m70 P82m71 p82w72 p82w73 p82w74 p82w75 p82w76 p82w77 p82w77		A A CT ACG CT ACG	CCT CCT CCT CT CCT CCT	TTC GTC GTC GTC GTC	AAT AAT AA AA AAC		ATT ATT ATT	G G GG	15 15 16 16 16 14 14	297 298 299 300 301 302 303 304 305
p82w79 p82w80 p82w81 p82w82 p82w83 p82w84 p82w85 p82w86 p82w87 p82w88	A CC	T ACG T ACA T ACA CT ACA A ACT ACA ACA A	CCT CCG CCG CCT CCT CCT	GTC GTC GTC GTC GTC GTC GTC	A AAC AA AAC AAC AAC AAC AAC	ATA ATA A AT	ATT	G	14 14 13 15 15 15 14 17	306 307 308 309 310 311 312 313 314 315
p82w88 p82w89 p82w90 p82w91 p82w92 p82w93 p82w94 p82w95 P82w96 p82w97 p82w98	CC	ACA TCA ACA CA	CCT CCT CCT CCT CCT CCT CCT	GTT GTC GTC GTC GTC GTC GTC	AAC AAC AAC AAC AAC AAC AAC AAC	ATA ATA ATA ATA ATA ATA	AG AT ATT ATT ATT	GG	15 16 14 16 15 16 16 14 15	316 317 318 319 320 321 322 323 324 325

p82m99		ACA	CCT	TTC	AAC	ATA	Α			16	326
p82m100	${f T}$	ACA	CCT	TTC	AAC	ATA				16	327
p82m101		ACA	CCT	ATC	AAC	ATA	ATG			17	328
P82m102		ACA	CCT	ATC	AAC	ATA	AG			16	329
p82m103		CA	CCT	GCC	AAT	ATA	ATG			16	330
p82m104			CCT	GCC	AAT	ATA	AG			16	331
p82m105			CCC							15	332
p82m106			CCC				AG			15	333
p82m107	Т	ACG			AAC					15	334
p82w108	CT		CCG				:			14	335
p82w109	·CCT						•			14	336
p82w110			CCG			ATA	ATG			15	337
p82w111			CCG							16	338
p82w112	СТ		CCA							14	339
p82w113	CT		CCA.	-		Α				15	340
p82w114			CCA							15	341
p82w115			CCA				AG			16	342
p82w116	. Т		CCT							15	343
p82w117	_		CCT							15	344
p82w118	т		CCT							14	345
p82m119	CCT		CCT		AAC					15	346
p82m120	CT		CCT		AAC					14	347
p82m121	A CCT	ACA		TTC	AA					15	348
p82w122			CCT	GTC	AAC	ATA	AGG			16	349
p82w123	Т	ACG	CCT	GTC	AAC	ATA				16	350
p82w124		CG	CCT	GTC	AAC	ATA	AGG			15	351
p82m125	Т	ACA	CCT	TTC	AAC	GTA				16	352
p82m126		ACA	CCT	TTC	AAC	GTA	AGG			16	353
p82m127			CCT		AAC		ATG			16	354
p82m128			CCT		AAC					16	355
p82o129					AAC			GGA	AGA	16	356
p82o130				C	AAC	GTA	ATT	GGA	AG	15	357

	86		88	89	90	91	92	93	94	length	Seq ID
	GGA	AGA				ACT		ATT	GGT		
P90w1		Α	-		TTG		CAG			16	358
P90w2		GA	AAT	CTG	TTG		CAG			17	359
P90w3		GA	AAT	CTG	TTG	ACT	CAG	AGG		18.	360
P90w4		Α	AAT	CTG	TTG	ACT	CAG	AGG		· · 17	361
P90w5		AGA			TTG		CAG			19	362
P90w6		AGA	AAT	CTG	TTG	ACT	CAG	ATG		20	363
P90w7		AGA	AAT	CTG	TTG	ACT	CAG	ATT		21	364
P90w8	AC	SA AA	T CI	rg Ti	G, AC	CT CF	AG A	TGG		20	365
P90w9	GA	AGA	AAT	CTG	TTG	ACT	CAG	AGG		21	366
P90w10	Α	AGA					CAG	ATG		21	367
P90m11		AGA	AAT	CTG	ATG	ACT	CAG			20	368
P90m12		AGA	AAT	CTG	ATG	ACT	CAG	ATT		21	369
P90m13	Α	AGA	AAT	CTG	ATG	ACT	CAG	AGG		20	370
P90m14	GA	AGA	AAT	CTG	ATG	ACT	CAG	AGG		21	371
P90m15	Α	AGA	TAA	CTG	ATG	ACT	CAG	ATG		21	372
P90m16		AGA								20	373
P90m17	GGA	AGA	AAT	CTG	ATG	ACT	CAG			21	374
P90m18	Α	AGA	AAT		ATG		CAG			19	375
P90m19		Α	AAT			ACT			GG-	21	376
P90m20		Α	AAT	CTG	ATG	ACT	CAG	ATT	G	20	377
P90m21			AAT	CTG	ATG	ACT	CAG	CTT	G	20	378
P90m22		Α	AAT			ACT		CTT	•	19	379
P90m23			AAT	CTG	ATG	ACT	CAG	CTT	G	18 "	380
P90w24						ACT			G	20	381
P90w25		Α	AAT			ACT				19	382
P90w26			AAT	CTG	TTG	ACT	CAG	CTT	G	19	383
P90w27			AAT			ACT				14	384
P90w28			AAT	CTG	TTG		CAG			15	385
P90w29			AAT	CTG		ACT				15	386
P90w30		Α	AAT	CTG		ACT	CAG			16	387
P90m31			AAT		ATG		CA			14	388
P90m32			AAT			ACT	CAG			15	389
P90m33			AAT			ACT				15	390
P90m34			AAT			ACT				16	391
P90w35			AAT							15	. 392
P90w36		GA	ACT							15	393
P90w37			T			ACT		ATG		15	394
P90w38						ACT				15	395
P90w39		GA	ACT	CTG	TTG	ACT	C			15	396
P90w40		Α				ACT				15	397
P90w41						ACT				15	- 398
P90m42			AAT			ACT				15	399
P90m43		A	AAT			ACT				15	400
P90w44			AT			ACT			•	15	401
P90w45						ACT		ATI		15	402
P90w46		AGA	AAT			ACT				15 15	403
P90m47			ΤA	CTG	ATG	ACT	CAG	AG		15	404

Table 3 - Cont'd-11

									•	
P90m48			CTG .	ATG	ACT	CAG	ATT		15	405
P90m49	AGA	AAT	CTG	ATG	ACT	CA	•		17	406
P90w50			ATG	TTG	ACT	CAG			15	407
					ACT		•		16	408
P90w51	GA				ACT				15	409
P90w52	07								16	410
P90w53	GA				ACT	CA				411
P90w54					ACC				15	
P90w55	Α				ACC				15	412
P90m56			ATG		ACC	CAG	:		15	413
P90m57	A	CAG	ATG	ATG	ACC	CA			15	414
P90w58		AAC	ATG	TTG	ACT	CAG			15	415
P90w59	A				ACT	CAG			15	416
P90w60					ACT	CAG	CTT		14	417
P90w61			CTG			CAG	CTG		14	418
					ACT	CAG	CTT		14	419
P90m62					ACT	CAG			14	420
P90m63						CAG			14	421
P90w64									14	422
P90w65			CTG			CAG	C-G		15	423
P90w66		AAT	CTG.		ACA					424
P90w67		AAC	CTG	TTG	ACT	CA			13	
.P90w68		AAC	CTG	TTG	ACT	С			13	425
P90w69	GA	AAC		TTG	ACT			_	13	426
p90w70			ΤG	TTG			ATT	G	15	427
p90w71			TG	TTG	ACT		ATT	GGG	16	428
p90w72			G	TTG	ACT		ATT	GGG	15	429
p90w73			TG	TTG		CAG		G	15	430
p90w74			CTG	TTG		CAG			15	431
p90w75			G	TTG	ACA	CAG	CTT	GGG	15	432
p90w76			TG	TTG	ACT	CAG	CTT	G	15	433
p90w77			G	TTG	ACT	CAG	ATG		15	434
p90w78			Ğ		ACT		CTT	G	14	435
p90w79			TG				ATT	G	15	436
p90w79			Ğ	TTG			ATT	G	14	437
			Ğ	TTG				GGG	15	438
p90w81			TG				ATT	G .	15	439
p90m82			TG		ACT			GGG		440
p90m83					ACT			GGG	15	441
p90m84			G G		ACT				16	442
p90m85								GGI	15	443
p90m86			CTG	ATG	ACI	CAG	CTT	<u></u>	15	444
p90m87	_						CTT	G		
P90w88		AAT			ACT				15	445
P90w89		AAT			ACT				15	446
p90w90	A	AAT					_		15	447
p90w100			CTG				;		15	448
p90m92	A	AAT	CTG	ATG	ACI	CA			16	. 449
p90m93	GA	AAT	CTG	ATC	ACI	. C			15	450
p90m94			CTG			CAG	ATG		15	451
p90m95	AGA	LAAT	ATG	ATO	;				15	452
p90m96	A AGA	LAAT	ATG	ATO	ACI				16	453
-										

0007	75	$\Lambda \subset \Lambda$	AAT	CTG	Δ Τ С	ACT				16	454
p90m97										16	455
p90m98	Α	AGA	AAT	ATA			030				456
p90m99		Α				ACT				16	
p90m100			AAT			ACC				15	457
p90m101			AAC	CTG	ATG	ACT	CAG			15	458
p90m102		AGA	AAT	TTG	ATG	ACT	C .			16	459
p90m103		A	AAT	TTG	ATG	ACT	ATG	ACT		16	460
p90m104				CTG		ACT	CAG			14	461
				CTG		ACT	CAG	A		16	462
p90m105			AT			ACT				16	463
p90m106			AT		ATG		CAG			14	464
p90m107			AI				CAG	Vulu	G	16	465
p90m108								VII	G	16	466
p90m109			AAT				C ·				
p90m110			AAT		ATG					15	467
p90m111	GA		\mathbf{AAT}	CTG	ATG	A				15	468
p90m112	GGA	AGA	AAT	CTG	ATG	A				16	469
p90m113	GA		AAT	CTG	ATG	AC				16	470
p90m114		AGA	AAT	CTG	ATG	AC				14	471
p90w115			AAT	CTG	TTA	ACT	CAG			15	472
p90w116			T	CTG		ACT		ATT		16	473
			ΑT				CAG			15	474
p90w117		7 (7		TTG					•	16	475
p90w118		AGA					C			15	476
p90w119		GA	_	TTG						15	477
p90w120			AAT	TTG	TTG	ACT	CAG			13	4//

Table 5

		non-B	86	7	4	9	0	
		Type B	95.7	1.1	8.5	1.1	1.1	
	probes tor	codon p50	w31	w44	w52	m3.7	neg.	
		non-B	70	22	4	0	œ	
		Type B	71.3	11.7	16	3.2	0	
	probes for	codon p48	W4.7	w45	w72	m41	neg.	
_		non-B	98	0	1	0	0	Н
		Type B	95.7	1.1	1.1	1.1	1.1	0
	probes for	codon p30 Type	w25	w29	w32	w36	m23	ned.

Table 5 - Cont'd

non-B 2.5 17.5 0 65 2.5 22.5 22.5 10 5 2.5 12.5
Type B 50 66.1 7.1 12.5 7.1 5.4 0 0 19.6 0
probes for w27 w37 w39 w50 w50 w69 w73 m69 m43 m56 neg.
non-B 70 12 12 0 0 0 0 0 0 0 0 2 2 2 2 2 2 2 3 3 8 8 8 8 8 8 8 8 8 8 8
Type B 81.9 2.1 1.1 1.1 2.1 2.1 1.1 1.1 1.1 1.3 3.2
probes for codon p82/84 w91 w60 w111 w89 w42 m36 m36 m67 m38 m105 m105 m107 m40 m63
18 62 62 18 0 0 4 4 4 4 4
1770e B 71.3 81.9 3.2 6.4 4.3 0 0 3.2 14.9
probes for codon p54 w3 w34 w14 w19 w22 w26 w27 m35 m35 neg.

Table 6

p30 ⁻	USA	France	U.K.	Brazil	Spain	Luxemb.	Belgium
w25	98.9	99.4	88.9	98.3	94.3	100.0	97.0
w29	2.5	0.6	0.0	1.7	0.0	0.0	0.0
w32	3.3	0.6	5.6	5.2	5.7	6.7	1.5
w36	2.5	0.0	0.0	. 3.4	0.0	0.0	1.0
m23	3.1	0.0	0.0	0.0	0.0	0.0	1.0
neg.	0.6	0.6	5.6	0.0	0.0	0.0	1.0
3							
p46/48	USA	France	U.K.	Brazil		Luxemb.	Belgium
w47	94.2	80.5	83.3	89.7			82.9
w45	8.6	15.6	0.0	1.7			11.1
w72	4.2	0.0	16.7	0.0	2.9		
m41	0.0	0.0	0.0	10.3	•		•
neg.	2.8	4.5	0.0	0.0	0.0	0.0	2.5
			•		_	_	
p50	USA	France	U.K.		_		_
w31	96.4	97.4	100.0	96.6	100.0	100.0	96.5
w44	1.7	0.6	0.0	1.7	0.0	0.0	1.0
w52	10.0	4.5	0.0	1.7	2.9		
m37	2.5	0.6	0.0	1.7		6.7	
neg.	3.1	2.6	0.0	3.4	0.0	0.0	1.5
p54	USA	France	U.K.	Brazil	Spain	Luxemb.	Belgium
w34	96.9	82.5	97.2	87.9	100.0	53.3	89.4
w3	84.7	77.9	94.4	69.0	82.9	46.7	76.9
w14	3.3	5.8	0.0	3.4	11.4	0.0	6.5
w19	9.2	2.6	0.0	1.7	2.9	6.7	5.5
w22	2.8	10.4	0.0	0.0	5.7	0.0	2.5
w26	0.0	1.3	0.0	0.0	0.0	0.0	0.0
w27	0.0	1.9	0.0	0.0	0.0	0.0	0.5
m55 .	0.0	0.0	0.0	0.0	0.0	13.3	0.5
m35	1.1	0.0	2.8	6.9	0.0	46.7	3.0
m37	0.0	0.0	0.0	0.0			
neg.	0.6	1.3	0.0	1.7	0.0	0.0	2.0
p82/84	USA	France	· u.ĸ.	Brazil	Spain	Luxemb.	Belgium
w91	91.6	93.5	94.4			73.3	85.9
w60	6.4		0.0			13.3	5.5
w111	3.6		0.0			0:0	0.5
w89	7.0		0.0			0.0	3.0
w42	0.6		2.8			0.0	2.0
. m36	0.3		0.0			0.0	0.0
m67	0.0		0.0			0.0	0.5

Table 6 - Cont'd

m38	0.0	0.0	0.0	0.0	0.0	6.7	0.0
	0.0	0.0	0.0	0.0	0.0	0.0	0.0
m105		0.0	0.0	0.0	0.0	0.0	0.0
m127	0.0					46.7	0.0
m40	2.8	0.0	8.3	3.4	5.7		
m63	0.3	0.0	0.0	1.7	2.9	13.3	0.5
m101	1.9	4.5	0.0	3.4	0.0	6.7	4.0
neg.	2.5	3.9	0.0	13.8	0.0	6.7	,5 . 0
p90	USA	France	U.K.	Brazil	Spain	Belgium	
w27	51.1	45.5	34.3	. 47.7	52.8	25.7	
w37	91.9	73.4	80.0	81.8	88.9	55.2	
w39	0.0	0.0	0.0	0.0	0.0	2.9	
w50	2.6	23.8	2.9	13.6	11.1	21.9	
w52	8.4	11.2	5.7	6.8		4.8	
w69	5.2	1.4	5.7	2.3	0.0	3.8	
w73	6.1	9.1	0.0	0.0	8.3	6.7	
w79	7.1	11.2	8.6	9.1	5.6	5.7	
m43	1.9	0.0	11.4	0.0	0.0	8.6	
m56	0.3	1.4	0.0	0.0	0.0	0.0	
neg.	1.0	0.0	0.0	0.0	0.0	18.1	

Table 7														
									Tm	lengt	e Se	eq ID		
	AGG	GGG	AAT	TGG	AGĞ	TTT	TA				20		511	
							32 GTA							

pc50w5	AGG GGG AAT TGG AGG TTT TA	20	511
pc30w25 pc30w29 pc30w32 pc30w36 pc30m23	26 27 28 29 30 31 32 33 34 35 ACA GGA GCA GAT GAT ACA GTA TTA GAA GAA GCA GAT GAT ACA GT A GCG GAT GAT ACA GCA GAT GAC ACA GT GCA GAC GAT ACA GT 40 A GCA GAC GAT ACA GT 40 A GCA GAT AAT ACA GT 40	14 13 14 14	31 35 38 42 29
pc48w37 pc48w47 pc48w73 pc48w45 pc48w72 pc48m41	44 45 46 47 48 49 50 51 52 CCA AAA ATG ATA GGG GGA ATT GGA GGT	15 15 14 18 16 15	512 93 513 91 120 87
pc50w31 pc50w44 pc50w52 pc50m37	48	15 14 14 12	151 164 172 157
pc54w34 pc54w14 pc54w19 pc54w22 pc54w26 pc54w27 pc54m35 pc54m37 pc54m55	51 52 53 54 55 56 57 58 GGA GGT TTT ATC AAA GTA AGA CAG GA GGT TTT ATC AAA GTA 42 A GGC TTT ATC AAA GTA 42 GA GGT TTT ATT AAA GTA 42 A GGT TTT ATT AAG GTA 42 GGT TTT ATT AAG GTA 40 GGT TTT GTC AAA GTA 40 GGT TTT GCC AAA GTA 42 A GGT TTT GCC AAA GTA 42	16 16 17 16 16 15 15	212 189 194 197 202 204 213 215 516
pc82w91 pc82w60 pc82w111 pc82w89 pc82m101 pc82w42 pc82m38 pc82m105 pc82m127	78 79 80 81 82 83 84 85 86 87 GGA CCT ACA CCT GTC AAC ATA ATT GGA AGA ACA CCT GTC AAC ATA A CA CCT GTC AAT ATA ATG A CCG GTC AAC ATA ATT ACA CCT GTT AAC ATA ATT ACA CCT GTT AAC ATA ATT ACA CCT ATC AAC ATA AT CA CCT GTC AAC ATA AT ACA CCT TTC AAC GTA ACG CCC TTC AAC ATA ACG CCC TTC AAC GTA ATG ACG CCC TTC AAC GTA ATG	16 17 16 17 17 14 15 15	318 287 338 316 517 269 265 332 354

pc82m40 pc82m63 pc82m36 pc82m67			C <i>I</i> AC <i>I</i>	A CCI	GCC	CAA C	C ATA C ATA C GT	A AG			44 42	15 16 15 14	267 290 518 519
	86	87	88	89	90	91	92	93	94				
· ·	GGA	AGA	AAT	CTG	TTG	ACT	CAG	ATT	GGT				•
pc90w27			AAT	CTG	TTG	ACT	CA				38	14	384
pc90w37			T	CTG	TTG	ACT	CAG	AT				15	514
pc90w39		GA	GTC	AAC	AGA	GTT	C					15	515
pc90w50			AAT	ATG	TTG	ACT	CAG				40	15	407
pc90w52			AAT	TTG	TTG	ACT	CAG				40	15	409
pc90w69		GA	AAC		TTG						40	14	426
pc90w73				TG	TTG	ACA	CAG	CTT	G	•	44	15	430
pc90w79				TG	TTG	ACC	CAG	ATT	G		44	15	436
pc90m138	•	GT	CAT	C AG	A TT	r ct						14	510
pc90m56			AAT	ATG	ATG	ACC	CAG				42	15	413

15

35

CLAIMS

- 5 1. Method for determining the susceptibility to antiviral drugs of HIV viruses in a biological sample, with said method comprising:
 - a) if need be, releasing, isolating or concentrating the polynucleic acids present in the sample;
 - b) if need be amplifying the relevant part of the protease gene of HIV with at least one suitable primer pair;
 - c) hybridizing the polynucleic acids of step a) or b) with at least one of the following probes:

probes specifically hybridizing to a target sequence comprising codon 30; probes specifically hybridizing to a target sequence comprising codon 46 and/or 48; probes specifically hybridizing to a target sequence comprising codon 50; probes specifically hybridizing to a target sequence comprising codon 54; probes specifically hybridizing to a target sequence comprising codon 82 and/or 84; probes specifically hybridizing to a target sequence comprising codon 90; or the complement of said probes;

- further characterized in that said probes specifically hybridize to any of the target sequences presented in figure 1, or to the complement of said target sequences;

 d) inferring from the result of step c) whether or not a mutation giving rise to drug resistance is present in any of said target sequences.
- 25 2. Method according to claim 1, further characterized in that said polynucleic acids of step a) or b) hybridize with at least two of the said probes, or to the complement of said probes.
- 3. Method according to claim 2, further characterized in that said probes are chosen from the following list: seq id no 7 to seq id no 477, seq id no 510 to seq id no 519 or the complement of said probes.
 - 4. Method according to any of claims 1 to 3, further characterized in that said primer pair is chosen from the following primers: seq id no 3, seq id no 503, seq id no 504, seq id no 4, seq id no 506, seq id no 507, seq id no 508 and seq id no 509.
 - 5. Method according to any of claims 1 to 3, further characterized in that:

10

25

step b) comprises amplifying a fragment of the protease gene with at least one 5'-primer specifically hybridizing to a target sequence located at nucleotide position 210 to 260 of the protease gene, in combination with at least one suitable 3'-primer, and step c) comprises hybridizing the polynucleic acids of step a) or b) with at least one of the probes specifically hybridizing to a target sequence or its complement, comprising codon 90.

- 6. Method according to any of claims 1 to 3, further characterized in that:

 step b) comprises amplifying a fragment of the protease gene with at least one 3'-primer specifically hybridizing to a target sequence located at nucleotide position 253 (codon 85) to position 300, in combination with at least one suitable 5'-primer, and step c) comprises hybridizing the polynucleic acids of step a) or b) with at least one of the probes specifically hybridizing to a target sequence or its complement, comprising any of codons 30, 46, 48, 50, 52, 54, 82 and 84.
- Method according to claim 5, further characterized in that the 5'-primer is seq id 5 and the 3'primer is one primer or a combination of primers chosen from the following primers: seq id no
 4, seq id no 506, seq id no 507, seq id no 508 and seq id no 509.
- Method according to claim 6, further characterized in that the 5'-primer is one primer or a combination of primers chosen form the following primers: seq id no 3, seq id no 503, seq id no 504 and the 3'-primer is seq id no 6.
 - 9. A probe as defined in any of claims 1 to 3, for use in a method for determining the susceptibility to antiviral drugs of HIV viruses in a biological sample.
- 10. A nucleic acid comprising a nucleotide sequence represented by any of the following SEQ ID numbers: SEQ ID NO 478, SEQ ID NO 479, SEQ ID NO 480, SEQ ID NO 481, SEQ ID NO 482, SEQ ID NO 483, SEQ ID NO 484, SEQ ID NO 485, SEQ ID NO 486, SEQ ID NO 487, SEQ ID NO 488, SEQ ID NO 489, SEQ ID NO 490, SEQ ID NO 491, SEQ ID NO 492, SEQ ID NO 493, SEQ ID NO 494, SEQ ID NO 495, SEQ ID NO 496, SEQ ID NO 497, SEQ ID NO 498, SEQ ID NO 499 and SEQ ID NO 500; or a fragment thereof, wherein said fragment consists of at least two contiguous nucleotides and contains at least one polymorphic nucleotide.
- 35 11. A primer as defined in any of claims 4 to 8, for use in a method for determining the susceptibility to antiviral drugs of HIV viruses in a biological sample.

- 12. A diagnostic kit enabling a method for determining the susceptibility to antiviral drugs of HIV viruses in a biological sample, with said kit comprising:
 - a) when appropriate, a means for releasing, isolating or concentrating the polynucleic acids present in said sample;
 - b) when appropriate, at least one of the primers of any of claims 4 to 6;
 - c) at least one of the probes of any of claims 1 to 3, possibly fixed to a solid support;
 - d) a hybridization buffer, or components necessary for producing said buffer;
 - e) a wash solution, or components necessary for producing said solution;
- f) when appropriate, a means for detecting the hybrids resulting from the preceding hybridization;
 - h) when appropriate, a means for attaching said probe to a solid support.

1/21

Figure 1

Codon 30

Codon 46/48

Codon 50

Codon 54

2/21

Figure 1 - Cont'd

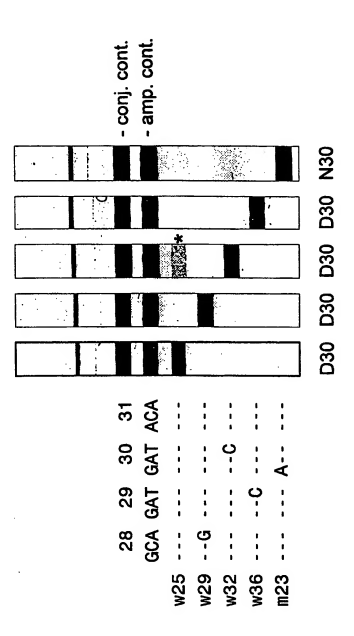
Codon 82/84

78	79	80	81	82	83	84	85	86	87
GGA	CCT	ACA	CCT	GTC	AAC	ATA		GGA	AGA
	A	T	G		С		G		
	G	T	С	\mathbf{A}_{i}	T		G		
		G	A	C			GG		•
				T			C		
				AC		•			
				TC	•				

Codon 90

·8 <i>6</i>	87	88	89	90	91	92	93	94
GGA	AGA	AAT	CTG	TTG	ACT	CAG	ATT	GGT
	C	C	A	A :	С	A	C	G
	A	С	T	C	A	A	G	C
	G		C	A	•		G	A
			A	AA			Ά	
			AA				GG	
							CG	





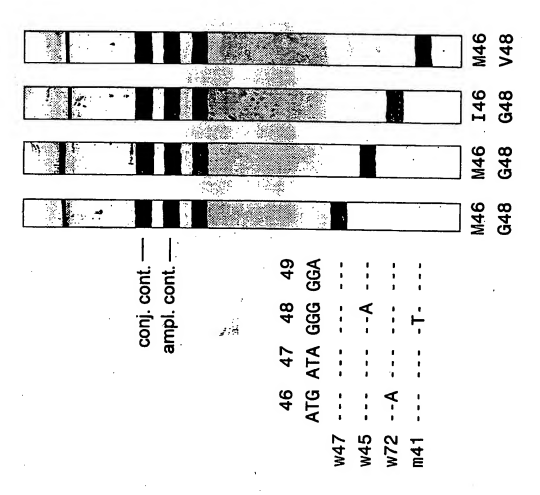


Figure 2B

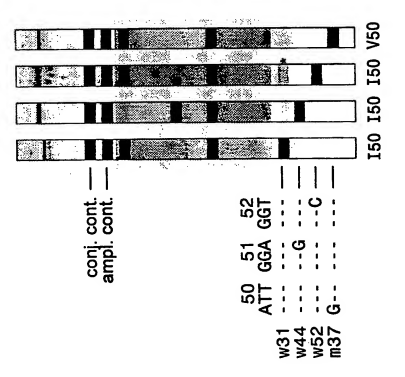


Figure 2C

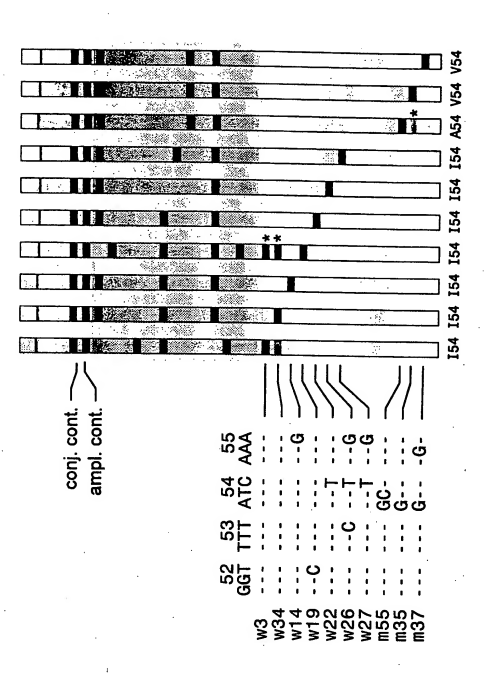
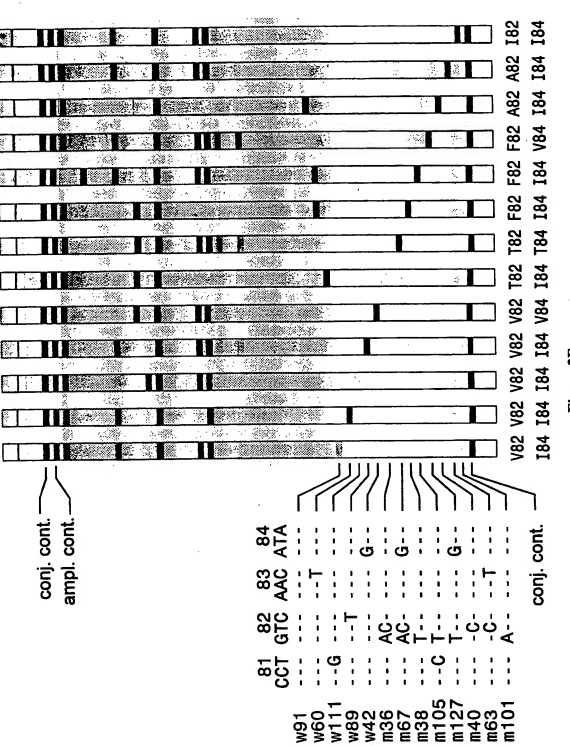


Figure 2I



ligure 2E

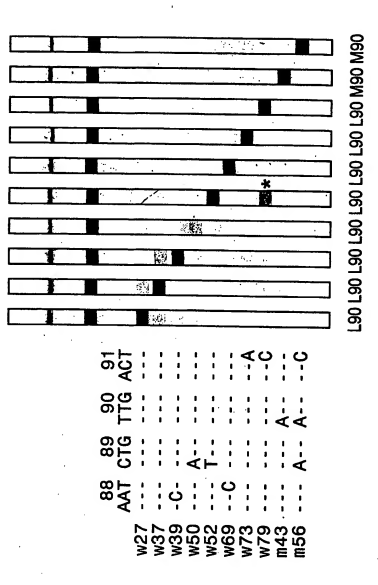
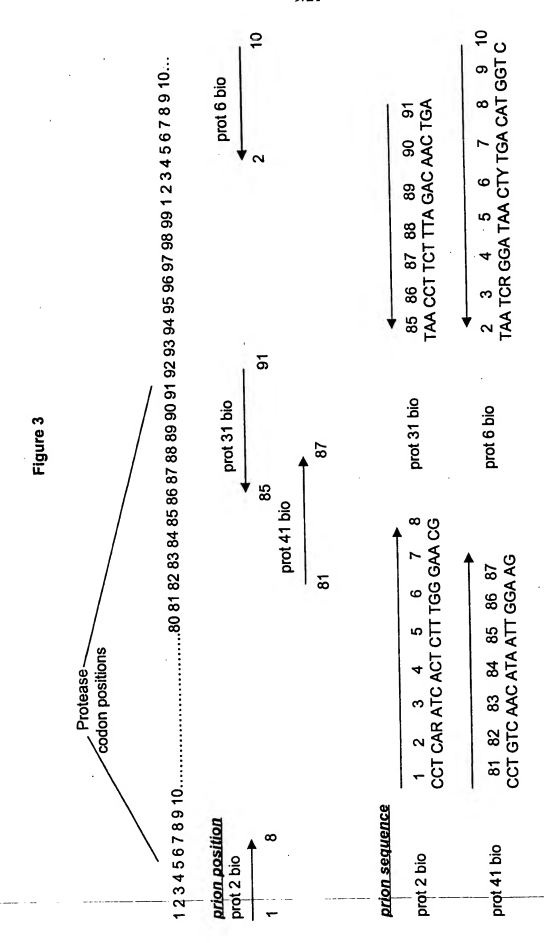
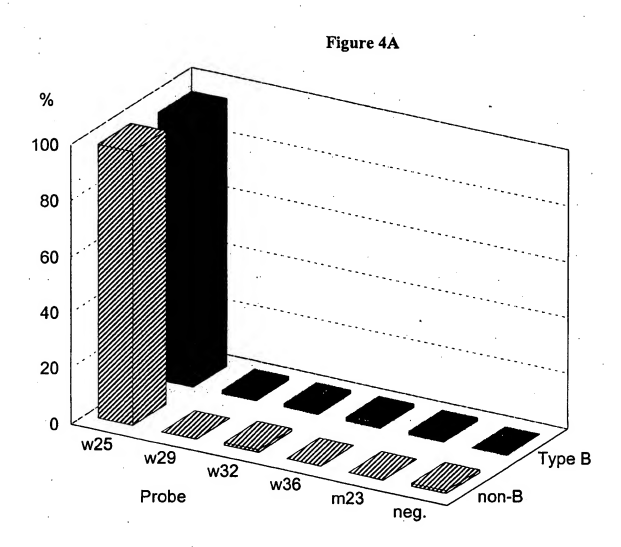


Figure 2F





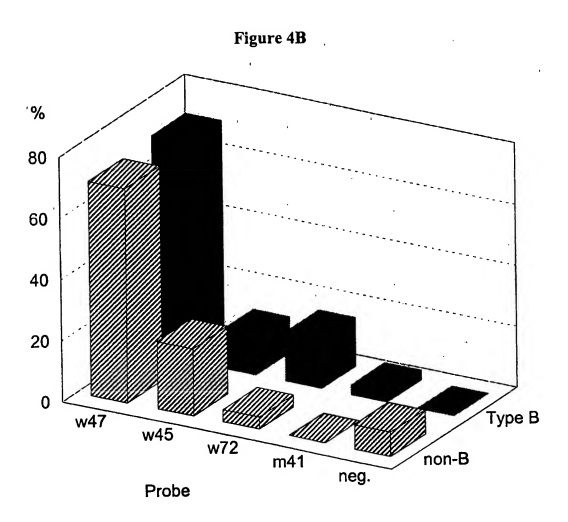
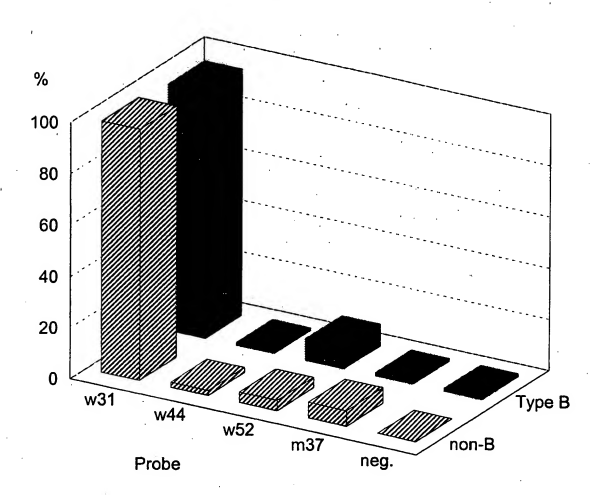
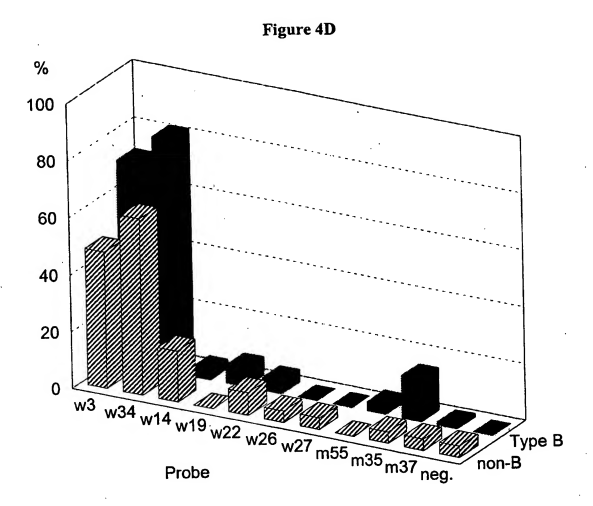
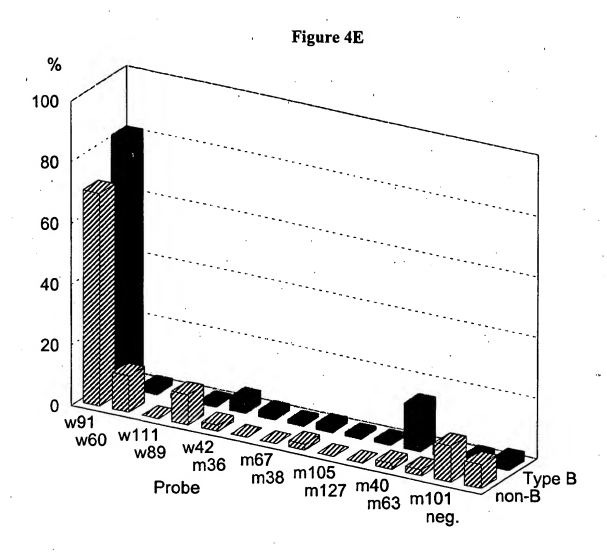


Figure 4C









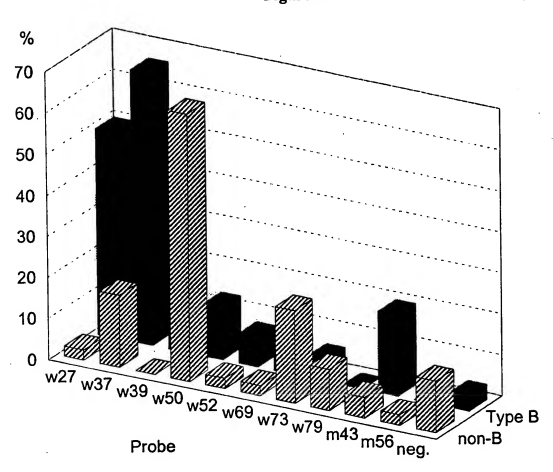


Figure 5A

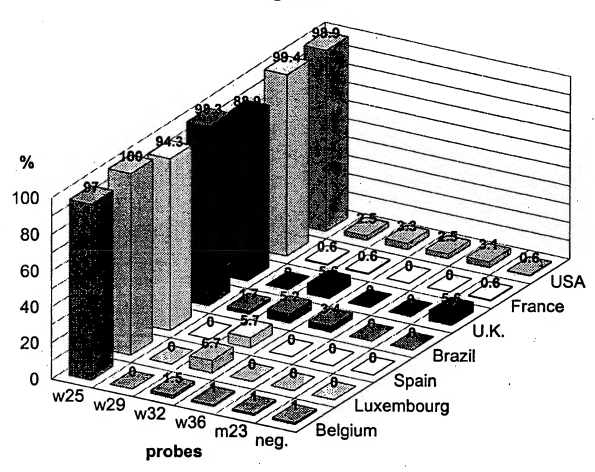


Figure 5B

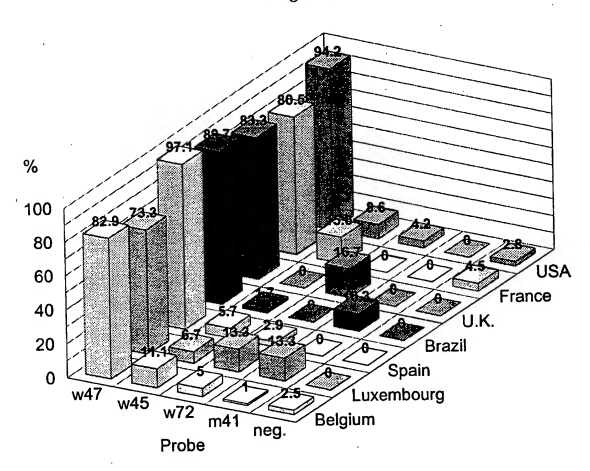
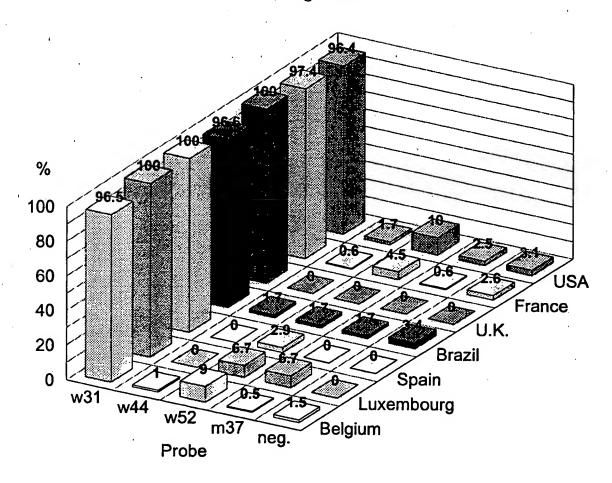
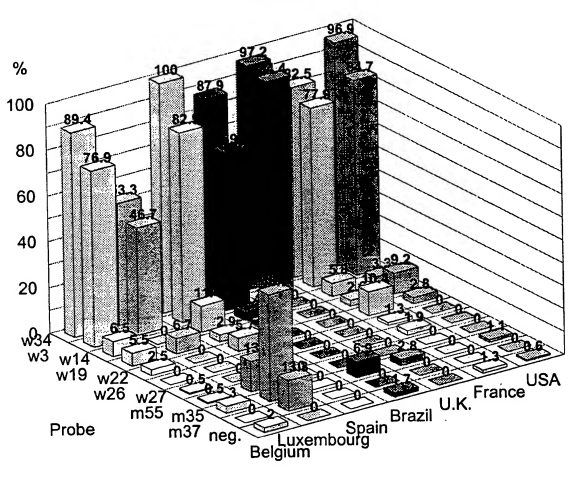
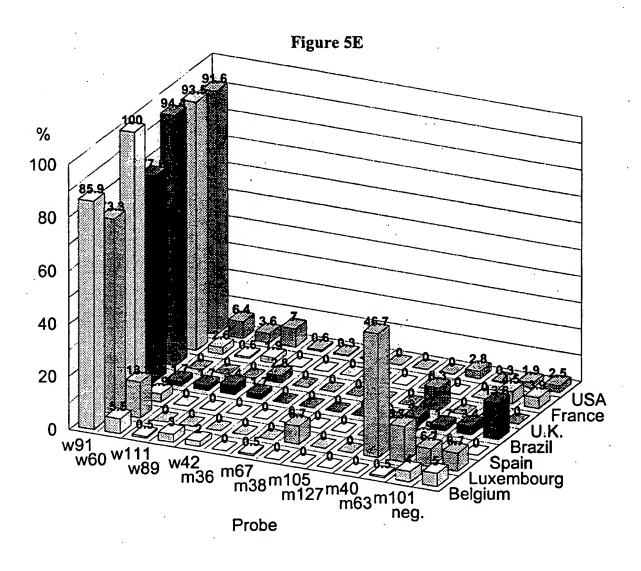


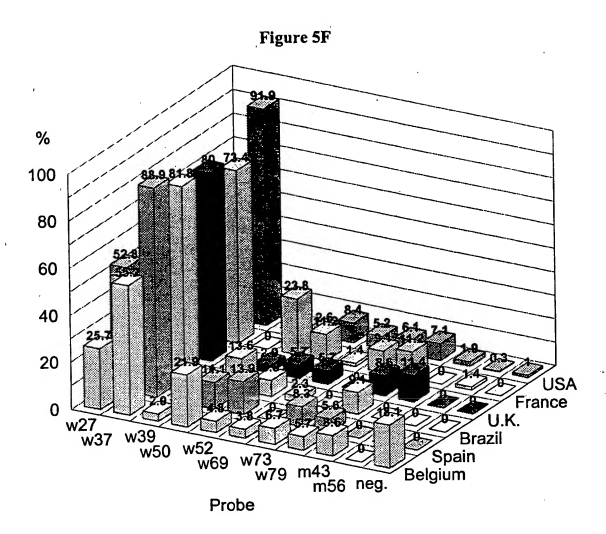
Figure 5C











PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT) (51) International Patent Classification 6: WO 99/67428 (11) International Publication Number: A3 C12Q 1/70 (43) International Publication Date: 29 December 1999 (29.12.99) PCT/EP99/04317 (21) International Application Number: (81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG. BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, (22) International Filing Date: 22 June 1999 (22.06.99) GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, (30) Priority Data: ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, 98870143.9 24 June 1998 (24.06.98) EP UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI (71) Applicant (for all designated States except US): INNOGENETpatent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, ICS N.V. [BE/BE]; Industriepark Zwijnaarde 7, P.O. Box 4, NE, SN, TD, TG). B-9052 Ghent (BE). (72) Inventor; and (75) Inventor/Applicant (for US only): STUYVER, Lieven Published [BE/BE]; Holestraat 8, B-9552 Herzele (BE). With international search report. (74) Common Representative: INNOGENETICS N.V.; Indus-

triepark Zwijnaarde 7, P.O. Box 4, B-9052 Ghent (BE).

(88) Date of publication of the international search report:

13 April 2000 (13.04.00)

(54) Title: METHOD FOR DETECTION OF DRUG-SELECTED MUTATIONS IN THE HIV PROTEASE GENE

(57) Abstract

The present invention relates to a method for the rapid and reliable detection of drug-selected mutations in the HIV protease gene allowing the simultaneous characterization of a range of codons involved in drug resistance using specific sets of probes optimized to function together in a reverse-hybridization assay. More particularly, the present invention relates to a method for determining the susceptibility to antiviral drugs of HIV viruses in a biological sample, with said method comprising: a) if need be, releasing, isolating or concentrating the polynucleic acids present in the sample; b) if need be amplifying the relevant part of the protease gene of HIV with at least one suitable primer pair, c) hybrydizing the polynucleic acids of step a) or b) with at least one of the following probes: probes specifically hybridizing to a target sequence comprising codon 30; probes specifically hybridizing to a target sequence comprising codon 46 and/or 48; probes specifically hybridizing to a target sequence comprising codon 54; probes specifically hybridizing to a target sequence comprising codon 50; probes specifically hybridizing to a target sequence comprising codon 50; probes specifically hybridizing to a target sequence comprising codon 90; or the complement of said probes; further characterized in that said probes specifically hybridize to any of the target sequences presented in figure (1), or the complement of said target sequences; d) inferring from the result of step c) whether or not a mutation giving rise to drug resistance is present in any of said target sequences.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

	•••						
AL AM AT AU AZ BA BB BE BF BG CF CG CH CI CM CN CU CZ DE DK EE	Albania Armenia Austria Austria Austriaia Azerbaijan Bosnia and Herzegovina Barbados Belgium Burkina Faso Bulgaria Benin Brazil Belarus Canada Central African Republic Congo Switzerland Côte d'Ivoire Cameroon Chima Cuba Czech Republic Germany Denmark Estonia	ES F1 FR GA GB GE GH GN HU IE IL IS IT JP KE KG KP KR LC LI LK LR	Spain Finland France Gabom United Kingdom Georgia Ghana Guinea Greece Hungary Ireland Israel Iceland Italy Japan Kenya Kyrgyzstan Democratic People's Republic of Korea Republic of Korea Republic of Korea Kazakstan Saint Lucia Liechtenstein Sri Lanka Liberia	LS LT LU LV MC MD MG MK ML MN MR MW MX NE NL NO NZ PL PT RO RU SD SE SG	Lesotho Lithuania Luxembourg Latvia Monaco Republic of Moldova Madagascar The former Yugoslav Republic of Macedonia Mali Mongolia Mauritania Malawi Mexico Niger Netherlands Norway New Zealand Poland Portugal Romania Russian Federation Sudan Sweden Singapore	SI SK SN SZ TD TG TJ TM TR TT UA UG US VN YU ZW	Slovenia Slovakia Senegal Swaziland Chad Togo Tajikistan Turkmenistan Turkey Trinidad and Tobago Ukraine Uganda United States of America Uzbekistan Viet Nam Yugoslavia Zimbabwe

In atlonal Application No PCT/EP 99/04317

A. CLASSIF	ICATION OF SUBJECT MATTER C1201/70		·
1100	01241/74		
	·		ĺ
According to	International Patent Classification (IPC) or to both national classif	ication and IPC	
B. FIELDS	SEARCHED cumentation searched (classification system followed by classification system followed by classification in the control of the cont	ation symbols)	
IPC 6	C12Q	2.00.09.000.0	
	• •		
Documentati	on searched other than minimum documentation to the extent tha	t such documents are included in the fields se	arched
Documentan	on searched other want manner coccurrence		
		, and the second second terms used	
Electronic da	ata base consulted during the international search (name of data	pase and, where practical, search terms occur	
			į
1	•		·
ł	·		
C. DOCUME	ENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.
X.	EASTMAN ET AL: "Genotypic chan		1-9
	human immunodefiency virus type	1	
	associated with loss of suppres plasma viral RNA levels in subj	sion of ects	
	treated with ritonavir (norvir)	-	•
	monotherapy"		
	JOURNAL OF VIROLOGY, June 1998 (1998-06),	
	XP002129272		
	the whole document		
		-/	•
			,
	·		
ļ			
1			
X Furt	ther documents are listed in the continuation of box C.	X Patent family members are listed	in annex.
° Special ca	ategories of cited documents:	"T" later document published after the into	ernational filing date
1 '	ent defining the general state of the art which is not	or priority date and not in conflict with cited to understand the principle or th	the application but
consi	dered to be of particular relevance document but published on or after the international	Invention "X" document of particular relevance; the	claimed invention
filing		carnot be considered novel or canno involve an inventive step when the do	t be considered to
which	ent which may liftow double on planty seamed or is cited to establish the publication date of another on or other special reason (as specified)	"Y" document of particular relevance; the	claimed invention eventive step when the
"O" docum	nent referring to an oral disclosure, use, exhibition or	document is combined with one or m ments, such combination being obvio	ore other such docu-
"P" docum	means nent published prior to the international filing date but	in the art. "&" document member of the same patent	
later	than the priority date claimed	Date of mailing of the international se	
Date of the	actual completion of the international search	Daily or manager	•
3	31 January 2000	11/02/2000	
Name and	mailing address of the ISA	Authorized officer	•
	European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk		
	Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Reuter, U	

Is ational Application No
PCT/EP 99/04317

	AND DESCRIPTION OF THE PROPERTY OF THE PROPERT	
C.(Continua Category	ation) DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Calegory		
X .	LIPSHUTZ R J ET AL: "USING OLIGONUCLEOTIDE PROBE ARRAYS TO ACCESS GENETIC DIVERSITY" BIOTECHNIQUES, US, EATON PUBLISHING, NATICK, vol. 19, no. 3, 1 September 1995 (1995-09-01), pages 442-447, XP000541924 ISSN: 0736-6205 the whole document	1,2,9
X	WO 97 41259 A (LACROIX JEAN MICHEL ;HUI MAY (CA); DUNN JAMES M (CA); LEUSHNER JAM) 6 November 1997 (1997-11-06) example 15	11
Y	CORDOBA J. ET AL: "'Human immunodeficiency virus and resistance!. VIRUS DE LA INMUNODEFICIENCIA HUMANA Y RESISTENCIAS." REVISTA ESPANOLA DE QUIMIOTERAPIA, (1998) 11/2 (152-156)., XP000867234 the whole document	1-9
Y	SCHINAZI ET AL: "Mutations in retroviral genes associated with drug resistance" INTERNATIONAL ANTIVIRAL NEWS, vol. 5, no. 8, August 1997 (1997-08), pages 129-142, XP000861634 cited in the application the whole document	1-9
Α	WO 97 27332 A (INNOGENETICS NV; STUYVER LIEVEN (BE); LOUWAGIE JOOST (BE); ROSSAU) 31 July 1997 (1997-07-31) the whole document	1-12
A	WINTERS ET AL: "Human immunodefiency virus type 1 protease genotypes and in vitro protease inhibitor susceptibilities of isolates from individuals who where switched to other protease inhibitors after long-term sequinavir treatment" JOURNAL OF VIROLOGY, vol. 22, no. 6, June 1998 (1998-06), pages 5303-5306, XP002129273 the whole document	1-12
	-/	

Ir. attonal Application No
PCT/EP 99/04317

C (Continue	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	/EF 99/0431/
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
P,X	SCHOOLMEESTER, A. (1) ET AL: "A line probe assay (LiPA) for the detection of drug-selected mutations in the HIV -1 protease gene." ABSTRACTS OF THE INTERSCIENCE CONFERENCE ON ANTIMICROBIAL AGENTS AND CHEMOTHERAPY, (1998) VOL. 38, PP. 396-397. MEETING INFO.: 38TH INTERSCIENCE CONFERENCE ON ANTIMICROBIAL AGENTS AND CHEMOTHERAPY SAN DIEGO, CALIFORNIA, USA SEPTEMBER 24-27, 1998 AMER, XP000869787 abstract	1-12
	•	
		•
	·	
	·	
	* ·.	
	•	
	·	

international application No.

PCT/EP 99/04317

- 1 Ob	ons where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This International Sea	arch Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.	: y relate to subject matter not required to be searched by this Authority, namely:
	.: 9, 10 and 12 by relate to parts of the International Application that do not comply with the prescribed requirements to such that no meaningful International Search can be carried out, specifically: RTHER INFORMATION sheet PCT/ISA/210
see rui	ATRIER THE OWNER, OF THE OWNER, OWNER
3. Claims Not because the	s.: ley are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Boy II Observa	tions where unity of invention is lacking (Continuation of item 2 of first sheet)
	earching Authority found multiple inventions in this international application, as follows:
This International S	earching Authority loans manipulation
As all required searchables	uired additional search fees were timely paid by the applicant, this International Search Report covers at e claims.
2. As all sea of any ad	urchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment ditional fee.
3. As only s	ome of the required additional search fees were timely paid by the applicant, this International Search Report nly those claims for which fees were paid, specifically claims Nos.:
4. No requirestricted	red additional search fees were timely paid by the applicant. Consequently, this International Search Report is It to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Prot	The additional search tees were accompanied by the applicant's protest.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.2

Claims Nos.: 9,10,12

Present claim 9 relates to a vast amount of nucleic acids so that a lack of conciseness within the meaning of Article 6 PCT arises to such an extent as to render a meaningful search of the claim impossible. Consequently the claimed nucleic acid sequences have not been searched per se.

Present claim 10 relates to an extremely large number of possible nucleic acid sequences so that a lack of clarity and conciseness within the meaning of Article 6 PCT arises to such an extent as to render a meaningful search of the claim impossible. Consequently, the search has been carried out for those parts of the claim which do appear to be clear and concise, namely the nucleic acid sequences themselves, which are specified with a sequence ID number. Neither nucleic acids comprising these sequences nor fragments of these sequences, wherin said fragment consists of at least two contiguous nucleotides and contains at least one polymorphic nucleotide, have been searched.

Present claim 12 relates to a vast amount of nucleic acids that a lack of clarity and conciseness within the meaning of Article 6 PCT arises to such an extent as to render a meaningful search of the claim impossible. Consequently the nucleic acid sequences being part of the claimed kit have not been searched per se.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

information on patent family members

Ir ational Application No PCT/EP 99/04317

Patent document cited in search report		Publication date	Patent family member(S)		Publication date
WO 9741259	Α	06-11-1997	US	5789168 A	04-08-1998
NO 3/41233	**	00 22 300.	US	588 8736 A	30-03-1999
			US	5830657 A	03-11-1998
			AU	2378097 A	19-11-1997
			AU	2747597 A	19-11-1997
			AU	2816797 A	19-11-1997
			AU	2816897 A	19-11-1997
			CA	2252487 A	06-11-1997
			CA	2252571 A	06-11-1997
			CA	2252588 A	06-11-1997
			WO	9740939 A	06-11-1997
			EΡ	0896632 A	17-02-1999
•			EP	0907752 A	14-04-1999
			EP	0914468 A	12-05-1999
			WO	9741257 A	06-11-1997
•			WO	9741258 A	06-11-1998
			US	5897842 A	27-04-1999
WO 9727332	Α	31-07-1997	AU	1444397 A	20-08-1997
WU JILIJJL	H	01 07 1557	BR	9704637 A	09-06-1998
		1	CA	2215073 A	31-07-1997
			EP	0817866 A	14-01-1998
			JP	11502727 T	09-03-1999

This Page is Inserted by IFW Indexing and Scanning Operations and is not part of the Official Record.

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images include but are not limited to the items checked:

BLACK BORDERS

IMAGE CUT OFF AT TOP, BOTTOM OR SIDES

FADED TEXT OR DRAWING

BLURRED OR ILLEGIBLE TEXT OR DRAWING

SKEWED/SLANTED IMAGES

COLOR OR BLACK AND WHITE PHOTOGRAPHS

GRAY SCALE DOCUMENTS

LINES OR MARKS ON ORIGINAL DOCUMENT

REFERENCE(S) OR EXHIBIT(S) SUBMITTED ARE POOR QUALITY

IMAGES ARE BEST AVAILABLE COPY.

As rescanning these documents will not correct the image problems checked, please do not report these problems to the IFW Image Problem Mailbox.

□ OTHER:

